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                 IPC reform
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        DEC 23 New IPC8 SEARCH, DISPLAY, and SELECT fields in USPATFULL/
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NEWS 9 JAN 13
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NEWS 17 FEB 22 The IPC thesaurus added to additional patent databases on STN
NEWS 18 FEB 22 Updates in EPFULL; IPC 8 enhancements added
NEWS 19 FEB 27 New STN AnaVist pricing effective March 1, 2006
NEWS EXPRESS FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
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              AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
              V8.0 AND V8.01 USERS CAN OBTAIN THE UPGRADE TO V8.01a AT
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=> file reg
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=>

Uploading C:\Program Files\Stnexp\Queries\10626012.str

chain nodes : 11 12 19 ring nodes : 1 2 3 4 5 6 7 8 9 10 13 14 15 16 17 18 chain bonds : 7-11 9-12 12-13 17-19 ring bonds : 1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10 13-14 13-18 14-15 15-16 16-17 17-18 exact/norm bonds : 7-11 16-17 17-18 17-19 exact bonds : 9-12 12-13 13-14 13-18 14-15 15-16 normalized bonds : 1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10 isolated ring systems : containing 1 : 13 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS 12:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS

L1 STRUCTURE UPLOADED

=> d l1 L1 HAS NO ANSWERS L1 STR

Structure attributes must be viewed using STN Express query preparation.

=> s l1 sample

SAMPLE SEARCH INITIATED 16:19:45 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

0 TO 0 0 0 TO 0 PROJECTED ITERATIONS:

PROJECTED ANSWERS:

L2 0 SEA SSS SAM L1

=> s l1 full

FULL SEARCH INITIATED 16:19:54 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED -2 TO ITERATE

100.0% PROCESSED 2 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

L3 0 SEA SSS FUL L1

=> s quinazol?

299314 QUINAZOL?

=> s hexahydro and pyrimid?

552247 HEXAHYDRO

1108000 PYRIMID?

L5 39615 HEXAHYDRO AND PYRIMID?

=> s 14 and 15

96 L4 AND L5

=> d scan 16

L6 96 ANSWERS REGISTRY COPYRIGHT 2006 ACS on STN
Asepino[2,1-b]quinasoline-3-carboxylic acid, 6,7,8,9,10,12-hexahydro12-oxo-, 2-(6-amino-1,2,3,4-tetrahydro-1,3-dimethyl-2,4-dioxo-5pyrimidinyl)-2-oxoethyl estar (9CI)
MF C22 H23 N5 06

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW HANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> file hcaplus COST IN U.S. DOLLARS

ENTRY SESSION 182.54 182.75

TOTAL

SINCE FILE

FULL ESTIMATED COST

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 16L72552 L6 => s 16/thu 2552 L6 755953 THU/RL L8 108 L6/THU (L6 (L) THU/RL)

96 S L4 AND L5

=> d his

L6

(FILE 'HOME' ENTERED AT 16:19:15 ON 28 FEB 2006)

FILE 'REGISTRY' ENTERED AT 16:19:24 ON 28 FEB 2006 L1STRUCTURE UPLOADED L2 0 S L1 SAMPLE L3 0 S L1 FULL L4 299314 S OUINAZOL? L5 39615 S HEXAHYDRO AND PYRIMID?

FILE 'HCAPLUS' ENTERED AT 16:21:39 ON 28 FEB 2006 L7 2552 S L6 L8108 S L6/THU

=> d l8 1- ibib abs fhitstr YOU HAVE REQUESTED DATA FROM 108 ANSWERS - CONTINUE? Y/(N):y

```
L8 ANSWER 1 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1335175 HCAPLUS
DOCUMENT NUMBER: 144:57603
Solid orally ingestible formulations of tetrodotoxin
INVENTOR(S): Lin, Weiyang
Can.
SOURCE: U.S. Pat. Appl. Publ., 14 pp.
CODEN: USXXCO
DOCUMENT TYPE: Patent
LANGUAGE: Patent
FAMILY ACC. NUM. COUNT: Patent INFORMATION:
PATENT NO. KIND DATE APPLICATION NO. DATE

US 2005222386 A1 20051222 US 2004-672528 20040622

WO 200512308 A1 20051229 WO 2005-CA973 20050621

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CH, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GH, HW, ID, IL, IN, IS, JP, KE, KG, FM, NP, KR, KZ, LC, LK, LK, LS, LT, LU, LV, MA, ND, NG, MK, MN, MY, MX, MZ, MA, NG, NI, NO, NZ, OM, PG, PH, FL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UM, GU, US, UZ, VC, VX, VX, VX, VZ, VZ, WZ, WZ, EF, ES, FI, FR, GB, GB, HU, IE, IS, IT, LT, LU, MC, NL, PL, FT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CH, GA, GN, GQ, GF, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO:

US 2004-872528 A 20046622

AB The present invention refers to outwardly solid or completely solid oral (or designed to be orally ingested) formulations of tetrodotoxin and/or analogs or derives. thereof.

IT 4368-28-9, Tetrodotoxin

RL: THU (Therapeutic use); PYP (Physical process); THU (Therapeutic use); BJOL (Biological study); PROC (Process); USES (USes)

(solid orally ingestible formulations of tetrodotoxin)
                                           (Uses)
(solid orally ingestible formulations of tetrodotoxin)
4368-28-9 HCAPLUS
5,9:7,10a-01methano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)
```

L8 ANSWER 2 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1293442 HCAPLUS
DOCUMENT NUMBER: 144:32262
INVENTOR(S): Spitzer, Nicholas C.; Borodinsky, Laurar Root, Cory M.
The Regents of the University of California, USA
PCT Int. Appl., 55 pp.
COUDENT TYPE: PANILY ACC. NUM. COUNT: 1 FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: ENT NO. KIND DATE APPLICATION NO. DATE

2005115367 A2 20051208 W0 2005-US16851
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BV, BY, BZ, CA, CR, CR, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GH, HR, HU, ID, IL, IN, IS, JP, KR, KG, KN, KP, KR, KZ, LG, LK, LK, LS, LT, LU, LV, MA, HD, MG, HK, MN, W, MX, MZ, AN, NG, NI, NO, NZ, CM, PG, PE, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SN, SY, TJ, TM, TM, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZW, ZW PATENT NO. WO 2005115367 NG, NI, No, NZ, CM, PG, PE, PT, RO, RU, SC, SD, SE, SG, SK, SL, SA, SY, TJ, TM, TM, TM, TM, TT, TZ, UA, UG, UZ, VC, VM, YU, ZA, ZM, ZW

RW BW, GR, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, TM, TM, TM, EB, BG, CH, CT, CZ, DE, DK, EE, ES, FI, FR, GB, GR, EU, IE, IS, IT, LT, LU, MC, NL, FL, FT, RO, SE, ST, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GR, GG, GW, ML, MR, NE, SS, TD, TG

PRIORITY APPLAN. INFO:

This application procvides, among others, a method for modulating the neurotransmitter activity of neurons, allowing for the treatment of various psychol. and neurol. disorders and illnesses. In one embodiment, a method of modulating neurotransmitter activity in a neuron associated with the central nervous system is provided. The method includes contacting the neuron with a stimulatory factor that alters the pattern of C22+ spike activity of the neuron. The neuron can be a fully differentiated adult neuron or embryonic neuron. The stimulatory factor can be elec. or chemical The neurotransmitter can be acetylcholine, nitric oxide, histamine, noradrenaline, a bioactive amine, an amino acid or a neuropeptide.

Generally, the modulation of neurotransmitter activity comprises altering neurotransmitter expression.

IT 4360-28-9, Tetrodotoxin
RN: FRC (Pharmacological activity); THU (Thexapoutic use); BIOL
(Biological study); USES (Uses)
(modulation of neurotransmitter activity in neurons by stimulatory factor that alters calcium spike activity for treatment of psychol. and neurol. disorders)

NA 4368-28-9 HCAPIUS

Absolute stereochemistry.

Absolute stereochemistry.

ANSWER 1 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

Absolute stereochemistry.

```
L8 ANSWER 3 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1050483 HCAPLUS
DOCUMENT NUMBER: 143:339667
TITLE: Compositions and methods to increase the effect of a neurotoxin treatment
                                                                                      David, Nathaniel E.
VYII NewCo 2003, Inc., USA
U.S. Pat. Appl. Publ., 13 pp.
CODEN: USXXCO
 INVENTOR(S):
 PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
                                                                                       Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 3
PATENT INFORMATION:
                                                                                                                                                        APPLICATION NO.
                 PATENT NO.
                                                                                      KIND
                                                                                                            DATE
                                                                                                                                                                                                                                        DATE
                            US 2005214325
                 WO 2005091891
            The present invention discloses compos. and methods for enhancing the effect (e.g., duration) of a neurotoxin treatment. The compos. herein include neurotoxins and neurotoxins and neurotoxins and neurotoxins conditions, urol. conditions, urol. conditions, urol. conditions, urol. conditions, urol. conditions, trol. compos. are administered locally to treat or prevent conditions, such as dermatol. conditions, urol. conditions, thyroid conditions, optical conditions, an eurol. conditions, tryled conditions, optical conditions, and neurol. conditions (Uses) (Uses) (Compos. and methods to increase effect of neurotoxin treatment) 4368-28-9 HCAPUS (Uses) (Compos. and methods to increase effect of neurotoxin treatment) 4368-28-9 HCAPUS (Spir) (John Compos. and methods to increase effect of neurotoxin treatment) (2-amino-104H-(1,3)dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-7,48,4aR,SR,75,95,10s,10aR,115,125)- (9CI) (CA INDEX NAME)
PRIORITY APPLN. INFO.
```

ANSWER 4 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
SSION NUMBER: 2005:647787 HCAPLUS
HENT NUMBER: 143:399552
E: Differential block of N-propyl derivatives of amitriptyline and dowepin for sciatic nerve block in DOCUMENT NUMBER: TITLE: rats
Gerner, Peter: Luo, Shi Huar Zhuang, Zhi-Yer Djalali,
Alimorad Gr. Zizza, Anthony M.; Myers, Robert R.;
Wang, Ging Kuo
Pain Research Center, Department of Anesthesiology,
Perioperative and Pain Medicine, Brigham and Women's
Hospital and Harvard Medical School, Boston, MA, USA
Regional Anesthesia and Pain Medicine (2005), 30(4),
344-350
CODEN: RAPMFX; ISSN: 1098-7339
Elsewier Inc.
Journal AUTHOR (5): CORPORATE SOURCE: SOURCE: DUBLISHER: CODEN: RAPMFK; ISSN: 1098-7339

PUBLISHER: Elsevier Inc.

DOCUMENT TYPE: Journal

LANGUAGE: English

AB The Pr group of ropivacaine (M-propyl-2',6'-pipecoloxylidide

hydrochloride) could be responsible for conferring some sensory

selectivity to this drug. Thus, adding a Pr group to exptl. local

anesthetics (LAs) (e.g., the tricyclic antidepressants amitriptyline and
doxepin) to increase sensory selectivity may be useful. We, therefore,

synthesized N-Pr amitriptyline and N-Pr doxepin and investigated a

potential predominance of sensory/nociceptive block over motor block

(differential block) in a rat sciatic nerve block model. In addition,
tetrodotomin (TTK), a naturally occurring Na* channel blocker, was

coinjected to investigate whether it increased block chiration. A 0.2-mL

test dose of N-Pr amitriptyline and N-Pr doxepin, at a concentration of 1,

2.5, 5, and 10 mM, (alone or in combination with TTX at a concentration of 20 was injected by the subfascial sciatic nerve approach. Motor function and sensory function (nociception) were evaluated by the force a rat's hind limb produced when pushing against a balance and the reaction to pinch, resp. N-Pr amitriptyline and N-Pr doxepin demonstrated prolonged block duration, with N-Pr anitriptyline displaying significant differential block at higher conces. (5 and 10 mM). The combination of either of these drugs with TTX increased the potency as well as the efficacy. Neurotoxicity commenced at concess of 5 to 10 mM. Detailed histopathol. nerve toxicity evaluations are justified to determine whether N-Pr amitriptyline has potential as a more sensory-selective local anesthetic at lower concess or as a predominantly sensory-selective neurolytic agent at higher concens.

4368-28-9, Tetrodotoxin
Ri: PAC (Pharmacological activity); TEW (Therepeutic use); BIOL (Biological study); USES (Uses)

(N-Pr amitriptyline, N-Pr domepin alone or in combination with tetrodotoxin demonstrated prolonged sciatic nerve block duration, N-Pr amitriptyline at higher dose displayed significant differential sciatic nerve block in rat)

4368-28-9 Bibliothano-10aH-[1,3]diomocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydromymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME) 5, and 10 mM, (alone or in combination with TTX at a concentration of 20 $\,$

ANSWER 3 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 4 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 5 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:592616 HCAPLUS
TITLE: Up-regulation of nNOS and associated increase in nitrecylc vasodilation in superior mesenteric arteries in pre-hepatic portal hypertension
AUTHOR(S): Jurzik, Lars; Froh, Matthlas; Straub, Rainer H.;
Scholmerich, Juercener, Wiest, Reiner

Jurzik, Lars: Froh, Matthias: Straub, Rainer H.: Scholmerich, Juergen Wiest, Reiner Department of Internal Medicine, University School of Medicine, Regensburg, 93042, Germany Journal of Repartology (2005), 43(2), 258-265 CODEN: JOREEC: ISSN: 0168-8278 CORPORATE SOURCE:

SOURCE:

PUBLISHER: Elsevier B.V. DOCUMENT TYPE:

LANGUAGE:

LISHER: Elsevier B.V.
UMENT TYPE: Journal
SUAGE: English
Splanchnic arterial vasodilation in portal hypertension has been
attributed largely to vascular NO overprodn. Three NO-synthase (NOS)
isoforms have been identified of which e(ndothelial)-NOS has been found
up-regulated and i (nducible)-NOS not expressed in the splanchnic
circulation in portal hypertension. So far, n(euronal)-NOS has not been
investigated and hence, the current study evaluates NNOS-expression and
nNOS-mediated vasorelaxation in a model of portal vein-ligated rats (PVL).
Mesenteric vasculature of PVL and sham rats was evaluated for nNOS-protein
(immunohistochem. and Western blotting). In vitro perfused
de-endothelialized mesenteric arterial vasculature was pre-constricted
with norepinephrine (ECBO) and tested for nNOS-mediated vasorelaxation by
periarterial nerve stimulation (PNS, 2-12 Rz, 45 V) before and after
incubation with the NOS-inhibitor L-NAME (10-4M). NNOS was localized to
the adventitia of the mesenteric arterial tree showing more intense
staining and increased protein expression in PVL as compared to sham rats.
PNS induced a frequency-dependent vasorelaxation, which was more
pronounced in PVL rats. L-NAME abolished this difference in
nerval-mediated vasorelaxation, the effect being significantly greater in
PVL than in sham animals. Perivascular antonS-protein expression is
enhanced in mesenteric arteries in portal hypertension mediating an
increased nerval No-mediated vasorelaxation. This nNOS-derived NO
overprodn. may play an important role in the pathogenesis of arterial
vasodilation in portal hypertension.
4368-28-9, Tetrodotoxin
RL BSU (Biological study, unclassified); TRU (Therapeutic use);
BIOL (Biological study); USES (Uses)
(tetrodotoxin completely blocked periatterial nerve stimulation induced
vasorelaxation in mesenteric artery in pre-hepatic portal hypertension
rat model)

act model)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,87,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 6 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:483312 HCAPLUS
DOCUMENT NUMBER: 143:188234
TITLE: Filtration and chromatograph for purifying

Filtration and chromatograph for puritying tetrodotoxin
Liang, Yinghua
Shanghai Huateng Bioengineering Co., Ltd., Peop. Rep.
China
Faming Zhuanli Shenqing Gongkai Shuomingshu, 7 pp.
CODEN: COXXEV INVENTOR(S): PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE PATENT NO. APPLICATION NO. DATE

PRIORITY APPLIA. INFO:

A 20030625 CN 2001-142658 20011214

PRIORITY APPLIA. INFO:: CN 2001-142658 20011214

AB Disclosed is a method for purifying tetrodotoxin from globefish. The method comprises cutting the viscers of globefish, milling, press filtering, deactivating to remove protein and grease, filtering through 1-5 µm filter membrane, 0.1-0.8 µm filter membrane, and then 1-5 nm filter membrane in sequence, purifying on chromatog. column, and crystallizing The purified tetrodotoxin may be used as sedative or analysis.

and is especially useful for reducing malignancy pain and drug withdrawal syndrome.

and is especially useful for reducing malignancy pain and use and is especially useful for reducing malignancy pain and useful syndrome.

A360-20-99, Tetrodotoxin
RL: BSU (Biological study, unclassified); PUR (Purification or recovery);
TMU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
(filtration and chromatograph for purifying tetrodotoxin for use as sedative and analgesic)
4360-28-9 HCAPLUS
5,917,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,105,10aR,115,125)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 5 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

REFERENCE COUNT:

THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 52

ANSWER 7 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:462110 HCAPLUS 143:76424 DOCUMENT NUMBER:

TITLE:

legs / 19324 Immunologic protection of anti-tetrodotoxin vaccines against lethal activities of oral tetrodotoxin

AUTHOR (5):

CORPORATE SOURCE:

Anny-Tal Beijing Institute of Pharmacology and Toxicology, Beijing, 100850, Peop. Rep. China International Immunopharmacology (2005), 5(7-8), 1213-1224 SOURCE:

CODEN: IINMBA; ISSN: 1567-5769

International Immunopharmacology (2005), 5(7-8), 1213-1224
CODEN: IINMAN, ISSN: 1567-5769
FUBLISHER: Elsevier B.V.
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Tetrodotoxin (TTX) is a high toxic small mol. neurotoxin. Haptenic vaccine for TTX was investigated and the carrier proteins were compared. TTX was conjugated to Tachypleus tridentatus hemocyanin (TTH) and tetanus toxoid (TT) via formaldehyde to form the artificial antigen TTX-TTH and TTX-TT. BALB/c mice were immunized with the artificial antigen TTX-TTH and TTX-TT. BALB/c mice were immunized with the artificial antigen the TTX-TTP. SALB/c mice were immunized with the artificial antigen the TTX-TTP. SALB/c mice were immunized with the artificial antigen the aice which exposed to TTX in dose of 600, 630, 800, 1220, 1500, 2000 and 2400 µg/kg survived at rates of 100, 100, 90, 90, 80, 50 and 201, with a LD50 value of 2020 µg/kg for TTH-TTX vaccine, and of 1001, 90, 91, 90, 91, 90, 91, 63, 61, 27.31 and 01, with a LD50 value of 1100 µg/kg for TT-TTX vaccine, resp. All control mice inoculated with carrier protein TTH or TT uniformly died of a dose of 600 µg/kg TTX i.g. challenge, half of them surviving about 6 mg/kg, and a few being able to bear a maximal accumulative dose as high as approx. 9 mg/kg of TTX challenges within eight months. The TTH-TTX vaccine was of the more excellent in protective effect from TTX oral intoxication, mainly resulted from the higher antibody affinity than that from TT-TTX vaccine vould high effectively protect animal from multiple, oral TTX intoxication.

IMMUNOPOPHYLAND ARCHIST STATE ARCHIST STATE OR STATE OR THE TX vaccine would high effectively protect animal from multiple, oral TTX intoxication.

IMMUNOPOPHYLAND ARCHIST STATE OR STATE

ANSWER 7 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 8 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 2005:369235 HCAPLUS MENT NUMBER: 142:404275 ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

Compositions and methods for enhancing cognitive function and synaptic plasticity Liu, Guosong Slutsky, Inna Massachusetts Institute of Technology, USA INVENTOR(S): PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 145 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English

	PATENT NO.					D	DATE			-	ICAT				D	ATE	
						-									-		
WO	2005	0372	15		A2		2005	0428		WO 2	004-	US33	971		2	0041	014
	¥:													BY,			
		CN,	co,	CR,	cυ,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	ΡI,	GB,	GD,
														KP,			
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA.	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		TJ,	TM,	TN,	TR,	TT,	TZ.	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZV
	RV:	BW,	GH,	GM,	ΧE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		ΑZ,	ΒY,	KG,	ΚZ,	MD,	RU,	ŦJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	ΗU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	BJ,	CF,	Œ,	CI,	CM,	GA,	GN,	GQ,	G₩,	ML,	MR,	NE,
		C11															

PRIORITY APPLN. INTO:

By the invention provides compns, and methods for enhancing cognitive function and synaptic plasticity. According to the method, Ca++ influx into excitatory neurons (nerve cells) is decreased by treatment with a number of the contraction of the contraction

of different agents including divalent cations (e.g., Mg++), GABAB agonists, GABAA agonists, calcium channel blockers, and/or compds. that decrease action potential fring such as sodium channel blockers. Decreasing Ca++ influx results in increased synaptic plasticity and enhanced cognitive function. In particular, decreasing Ca++ influx associated with uncorrelated neural activity results in long-lasting increases in synaptic plasticity and cognitive function. This is achieved by administration of agents that cause a voltage-dependent block of NMDA receptors (e.g., divalent cations such as Mg++) or by administration of GABAB agonists such as baclofen. The invention further provides screening methods useful in identifying compds. that enhance synaptic plasticity and cognitive function.

methods useful in identifying compds. that enhance synaptic plasticity cognitive function.
4368-28-9, TTX
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (compns. and methods for enhancing cognitive function and synaptic plasticity)
4368-28-9 HCAPBUS
5,917, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 9 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:322057 HCAPLUS
DOCUMENT NUMBER: 143:344497
Hyocytes from congenital myotonic dystrophy display abnormal Na+ channel activities
Bernareggi, Annalisa; Fucling, Denis; Mouly, Vincent; Ruzzier, Fablo; Sciancalepore, Marina
Department of Physiology and Pathology, University of Trieste, Trieste, 34127, Italy
FUBLISHER: COUNTY TYPE: John Wiley & Sons, Inc.
DOCUMENT TYPE: Journal

SOURCE:

Muscle & Nerve (2005), 31(4), 506-509

CODEN: MUNEDE, ISSN: 0148-639X

JOHN WILEY & Sons, Inc.

JOHN WILEY & MARCH &

REFERENCE COUNT:

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 10 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

HCAPLUS COPYRIGHT 2006 ACS on STN
2005:214987 HCAPLUS
143:264135
Sodium channels and neuropathic pain
Chung, Jin Mo: Chung, Kyungsoon
Department of Neuroscience & Cell Biology, University
of Texas Medical Branch, Galveston, TX, 77555-1069,

USA Novartis Foundation Symposium (2004), 261(Pathological Pain), 19-31 CODEN: NFSYF7, ISSN: 1528-2511 John Wiley & Sons Ltd. Journal; General Review SOURCE:

PUBLI SHER: DOCUMENT TYPE: LANGUAGE:

LISHER:
John Wiley & Sons Ltd.

MINDHT TYPE:
Journal: General Review
GUAGE:
English
A review. Although it has long been known that sodium channels play an important role in the generation of abnormal neuronal activity and neuropathic pain, it is only recently that we have begun to understand the subtypes of sodium channels which are particularly important in neuropathic pain. Many of the identified subtypes of sodium channels are localized in dorsal root ganglion (DMG) neurons. Based on their sensitivity to tetrodotoxin (TTX), these sodium channels are classified as TTX-sensitive (TTX) or TTX-resistant (TTX) subtypes. In in vitro electrophysiol. expts., ectopic discharges arising from DMG neurons with injured axons are blocked by TTX at doses that are too low to block TTX subtypes. These data suggest that TTXs subtypes of sodium channels are playing an important role in the generation of both ectopic discharges and neuropathic pain. Anal. of mRNA of the TTXs subtypes of sodium channels are playing an important role in the generation of both ectopic discharges and neuropathic pain. Anal. of mRNA of the TTXs subtypes of sodium channels are playing an important role in the generation of both ectopic discharges and neuropathic pain. Anal. of mRNA of the TTXs subtypes of sodium channels are playing an important role in the peneration of so the manufacture of the peneration of some paneration.

NER MR (MAG) are the only two subtypes that are up-regulated, suggesting their potentially important role in ectopic discharge and neuropathic pain (TTX subtypes of Na channels play role in generation of ectopic discharge and neuropathic pain, Nav.13 (Type III) and Nax (NaG) are two subtypes that are up-regulated suggesting their importance in pain generation in DRG neurons in rat)

1368-28-9 (TTX subtypes of Na channels play role in generation of ectopic discharge and neuropathic pain, Nav.13 (Type III) and Nax (NaG) are two subtypes that are up-regulated suggesting their importance in pain generation in DRG neurons in rat)

Absolute stereochemistry.

L8 ANSWER 11 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:85697 HCAPLUS
DOCUMENT NUMBER: 12:356973
ITILE: 12:356973
Drug-abstaining and analgesic medical formulation and its preparation
Lin, Wenhan
PATENT ASSIGNEE(S): 44 Penhan
Vang, Kaiye, Peop. Rep. China
Paming Zhuanli Shenqing Gongkai Shuomingshu, 7 pp.
CODDN: CNXEV
Patent
LANGUAGE: 45 Patent
Chinese
PATENT INFORMATION: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1485039	λ	20040331	CN 2002-131020	20020924
PRIORITY APPLN. INFO.:			CN 2002-131020	20020924
AB The drug-abstaining	and a	nalgesic inj	ection is composed of	tetrodotomin
0.1-20.0 μg, citrio	acid (0.5-100 pg,	and water 1 mL. Tetro	dotomin
is isolated by beat	ing ov	ary, viscus,	and skin of globefish	. vacuum

is isolated by overlang overlang overlang to the supernatant, dissolving the residual solid in 20% acetic acid solution, precipitating with ethanol, vacuum concentrating the supernatant to obtain

e tetrodotoxin, and purifying on C18 column.
4368-28-9, Tetrodotoxin
RL: TEU (Therapeutic use); BIOL (Biological study); USES (Uses)
(drug-abstaining and analgesic medical formulation and its preparation)
4368-28-9 BCAPUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,105,10aR,115,12s)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 10 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

3

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN 2005:64531 HCAPLUS 142:384963

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR (5): CORPORATE SOURCE:

142:384963
A microcapsule technique for long-term conduction block of the sciatic nerve by tetrodotoxin Martinov, Vladimir N., Nja, Arild Department of Physiology, Institute for Basic Medical Sciences, University of Oslo, Oslo, N-Osl7, Norway Journal of Neuroscience Methods (2005), 141(2), 199-205 SOURCE:

CODEN: JNMEDT: ISSN: 0165-0270 Elsevier B.V.

PUBLI SHER:

DOCUMENT TYPE:

ANGE: English
Tetrodotoxin (TTX) is a selective blocker of voltage-gated Na+ channels
that is used to block action potentials in vitro and in vivo. Maintaining
a sufficiently high local concentration of TTX in vivo to block conduction

peripheral nerve is tech. demanding and carries a risk of systemic toxicity. We report that slow diffusion of TTX out of a microcapsule (glass capillary) inserted beneath the epineurium of the sciatic nerve, with a loose cuff around the nerve, combines high blocking efficacy with low systemic toxicity in rats and mice. The local anesthesia and motor paralysis was stable for at least 4-6 wk. The conduction block was reversible and did not cause any obvious nerve injury. Low cost and simple surgical implementation make this new system an interesting alternative to existing long-term drug delivery methods.

4368-28-9. Tetrodotoxin
RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); TMU (Therapeutic use); BIOL (Biological study); USES (Uses)

activity); THO traerapeura use; BID (BIDIOGICAL SLODY), SUB-(Uses)

(tetrodotoxin via microcapsule technique inserted beneath epineurium of sciatic nerve reversibly blocked impulse conduction and did not cause nerve injury, showed low systemic toxicity in mouse and rat);
5,9:7,10a=Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 12 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 13 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 13 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:55070 HCAPLUS
142:141258
Stable tetrodotoxin freeze drying medicinal formulation containing disaccharides or polysaccharides as stabilizer
INVENTOR(5): Zhang, Xiao Kang, Yuhong; Huang, Xiaoyan PATENT ASSIGNEE(5): Rep. China PCT Int. Appl., 23 pp.
COURCE: PIXXD2

DOCUMENT TYPE: Patent DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent Chinese PATENT NO. APPLICATION NO. DATE KIND DATE WO 2004-CN736 A1 20050120 WO 2004-CN736 20040702

AM, AT, AU, AZ, BA, BB, BB, BR, BY, BY, BZ, CA, CH, CU, CZ, DE, DK, DM, DZ, EC, EE, EC, ES, FI, GB, GD, HH, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LT, LU, LV, MA, MD, MG, MX, MN, MY, MX, MZ, NA, NI, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TT, TZ, LA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, KE, LS, MY, MZ, MA, NI, SJ, SL, SZ, TZ, UG, ZA, ZY, ZY, MX, KZ, MB, RU, TE, TI, LU, MC, ML, PL, PT, RO, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, MS, 20040702 WO 2005004874 2005004874
Y: AE, AG, AL,
CN, CO, CR,
GE, GH, GM,
LK, LR, LS,
NO, NZ, CM,
TJ, TH, TN,
RW: BW, GH, GM,
AZ, BY, KG,
EE, ES, FI,
SI, SK, TR,
SN, TD, TG SN, TD, TG
CN 1568999 A 20050126 CN 2003-0146020 20030714
US 2005020610 Al 20050127 US 2004-890279 20040714
PRIORITY APPLN. INFO.: CN 2003-146020 A 20030714
AB Disclosure is a freeze drying preparation for injection containing in each dose 0.5 Disclosure is a freeze drying preparation for injection containing in each , 0.5 to 60kg tetrodotoxin or the analogs thereof, which has good stability and low toxicity, and can be stored at room temp for a long period of time. Said preparation also contains compds. which can reduce C-4 hydroxy activity of tetrodotoxin or the analogs thereof, such as glucosidic linkage containing compds. selected from any one of disaccharides, polysaccharide, the derivs. thereof or their mixture, and acid solubilizer which improves dissolving of tetrodotoxin or the analogs thereof. For example, an injection solution containing tetrodotoxin 3, lactose 3,000 (as stabilizer), citric acid 0.012 mg was frozen dried and showed improved stability comparing with fructose as stabilizer.
4368-28-9, Tetrodotoxin
RL: THU (Therepautic use); BIOL (Biological study); USES (Uses)
(tetrodotoxin freeze drying injections containing disaccharides or polysaccharides as stabilizers and acids as solubilizers for improved stability)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,10s,10aR,115,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 14 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:1045477 HCAPLUS
DOCUMENT NUMBER: 142:469133
Tetrodotoxin conjugate and its medical composition
XU, Qinhuir Rong, Kangtai
Institute of Toxic Medicine, Academy of Military
Medical Science of PLA, Peop. Rep. China
Faming Zhuanli Shenqing Gongkai Shuomingshu, 18 pp.
DOCUMENT TYPE: Patent
Patent
Patent
Patent DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

A 20040107 APPLICATION NO. DATE PATENT NO. KIND DATE

CN 1465403 A 20040107 CN 2002-123142 20020619

PRIORITY APPLN. INFO.:

CN 2002-123142 20020619

The conjugate of tetrodotoxin (TTX) with carrier (such as hemocyanin of limulus, tetanus toxold, or their fragments) is prepared by coupling TTX with hemocyanin (at a molar ratio of 250-7.0) in the presence of linker (1-24) such as formaldehyde or glutaraldehyde) at 30°C for 72 h, and then dialyling at 4°C to remove free toxin. The conjugate may be used to prepare monoclonal antibody, antiserum, or antitoxin as anti-TTX vaccine, also as immunol. affinity chromatog, reagent for purifying anti-TTX antibody, as immunol. affinity chromatog, reagent for purifying anti-TTX antibody, as immunoassay reagent for detecting TTX, further as the analytic reagent for studying electrophysiol. and pharmacokinetics of TTX, etc.

IT 4368-28-9, Tetrodotoxin
RL: PEP (Physical, engineering or chemical process); PTV (Physical process); TTV (Therexpeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (tetrodotoxin conjugate and its medical composition)

RN 4368-28-9 HCAPLUS

NS 59,77,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-mino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

L8 ANSWER 15 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:1022045 HCAPLUS
DOCUMENT NUMBER: 122:190938
AUTHOR(S): 122:190938
AUTHOR(S): Satoh, Elkin Hurskami, Keir Mishimura, Masakazu
Department of Pathobiological Science, Obihiro, Obihiro, Japan
SOURCE: International Journal of Neuroscience (2004), 114(5), 587-596
CODEN: INNUB7, ISSN: 0020-7454
Taylor 6 Francis, Inc.
DOCUMENT TYPE: JOURNAL

DOCUMENT TYPE:

LANGUAGE: English

In these studies, the authors investigated the effect of propylene glycol (PG) on the cytosolic free Ca2+ concentration ([Ca2+]i) in rat cerebrocortical

synaptosomes using the fluorescent Ca2+ indicator fura-2. PG (0.5-5% volume/volume) increased [Ca2+]i in a concentration-dependent manner. The

volume/volume) increased [Ca2+]i in a concentration-dependent manner. The nduced increase in [Ca2+]i seas inhibited approx. 50% by the omission of extracellular Ca2+ or the addition of Ni2+ (100 µM). Decrease of extracellular Na+ (6.2 mM) or addition of tetrodotoxin (1 µM), verapamil (10 µM), nifedipine (10 µM) or agatoxin IVA (200 mM), o-conotoxin GVIA (1 µM), or o-conotoxin MVIIC (1 µM) had no effect on the increase in [Ca2+]i. Also, addition of TMB-8 (100 µM), ryanodine (50 µM) or thappigargin (1 µM) did not modify the increase in [Ca2+]i in the absence of extracellular Ca2+. These results suggest that PG increases [Ca2+]i in rat cerebrocortical synaprosomes by both stimulating Ca2+ error TMB-8, ryanodine- and thappigargin-insensitive Ca2+ stores.

4368-28-9, Tetrodotoxin
RI: PAC (Pharmacological activity), TMU (Therapeutic use), BIOL (Biological study) USES (Uses)
[low Na+ in resting [Ca2+]i was higher than in presence of NaCl and voltage-dependent Na+ channel blocker tetrodotoxin had no effect in rat cerebrocortical synaprosomes

voltage-dependent has channel blocker tetrodotokin had no effect in cerebrocortical synaptosomes! 4368-28-9 HCAPUS 5,917,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-hydronymethyl)-,(4R,4aR,5R,75,95,105,10aR,11s,125)-[9CI] (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 16 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:969536 HCAPLUS
11TILE: 142:225714
Therapeutic agent for treatment of hemorrhoids using roe of globefish and production
LLIM, Gap Man
PATENT ASSIGNEE(S): S. Korea
SOURCE: RPWINT ASSIGNEE(S): S. Korea
CODEN: RWXXA7
DCCUMENT TYPE: Patent
LANGUAGE: KOREAN
KOREAN
CODEN: RWXXA7

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE

KR 2002064	907 A	20020810	KR 2001-5223	20010203
PRIORITY APPLN.	INFO.:		KR 2001-5223	20010203
AB A process	of preparing a	therapeutic	agent for hemorrhoids	by heating the

of a globefish at a specified temperature and then mixing sodium chloride is provided. The therapeutic agent for hemorrhoids is burned with alc. and a portion of hemorrhoids is exposed thereto. The roe of a globefish is heated at 0 to 30° for 50 to 150 days, ground and mixed with sodium chloride to produce a therapeutic agent for hemorrhoids containing tetrodotoxin.

4368-28-9. Therodotoxin
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(therapeutic agent for treatment of hemorrhoids using roe of globefish)
4368-28-9. HCAPLUS
5,917, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4,6,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 15 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 17 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 2004:943433 HCAPLUS HCAPLUS 142:204664

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

Pharmaceutical composition for treatment of cancer containing globefish extract Kim, Ik Soo S. Korea

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: Repub. Korean Kongkae Taeho Kongbo, No pp. given CODEN: KRXXA7

DOCUMENT TYPE: Patent

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
KR 2002091641	A	20021206	KR 2001-30491	20010531
PRIORITY APPLN. INFO.:			KR 2001-30491	20010531
Tetrodotoxin as a obtaining analgesi patient who suffer comprises a globef containing one or methotrexate, adri additive containin	main con c activi s from c ish ext more se amycin a g one of	mponent is painty while treatment paints and an allected from and taxol and a stabilized	ng a globefish extrac provided which has an eating cancer when ad . The pharmaceutical ictive ingredient, an the group consisting d a pharmaceutically er, favoring agent an	advantage of ministered to a composition anticancer agent of 5-fluorouracil, acceptable d corrigent. The

additive containing one of a stabilizer, favoring agent and corrigent. ovary of a globefish is extracted in water at 100°C for 24h and then centrifuged after removing suspended solids. 4368-28-9. Tetrodotoxin
RL: NPO (Natural product occurrence): THO (Therepeutic use):
BIOL (Biological study): OCCU (Occurrence): USES (Uses)
(antitumor compns. containing anticancer drugs and tetrodotoxin from globefish ovary as analgesics)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4,5,5,910-hexahydro-12-(hydroxymethyl)-(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

L8 ANSWER 18 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: HCAPLUS COPYRIGHT 2006 ACS on STN
2004:927018 HCAPLUS
141:388733
Compositions of a cyclooxygenase-2 selective inhibitor
and a sodium ion channel blocker for the treatment of
central nervous system damage
Stephenson, Diane T., Taylor, Duncan P.
Pharmacia Corporation, USA
PCT Int. Appl., 164 pp.
CODEN: PIXXOZ
Patent

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English 3 FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

ENT NO. KIND DATE APPLICATION NO. DATE

2004093811 A2 20041104 W0 2004-US12383 20040421
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, C2, DE, OK, DM, DZ, EC, EE, GC, ES, FI, GB, GD, GE, GH, GH, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LL, LW, LW, MA, MD, MG, MK, MM, MW, MX, MZ, NA, NI, NO, NZ, CM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, BW, GR, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, ST, SF, TP, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, HL, MR, NE, SN, TD, TG PATENT NO. WO 2004093811 A1 20041111 20040421

US 2004-829009 US 2003-464499P US 2003-464830P US 2004224940 PRIORITY APPLN. INFO.: P 20030422 P 20030423 OTHER SOURCE(S): MARPAT 141:388733

US 2003-64830P P 20030423

The invention provides compns. and methods for the treatment of central nervous system damage in a subject. More particularly, the invention provides a combination therapy for the treatment of a central nervous system ischemic condition or a central nervous system traumatic injury comprising the administration to a subject of a sodium ion channel blocker in combination with a cyclooxygenase-Z selective inhibitor. Use for the treatment of stroke is specifically claimed.

4368-2e-9, Tetrodotoxin

RL: PAC (Pharmacological activity): THU (Therapeutic use): BIOL (Biological study): USES (Uses) (cyclooxygenase 2 inhibitor-sodium channel blocker combination for treatment of CNS damage)

4368-28-9 HCAPUS

5,9:7,10a-Dimethano-10ai-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (48,4a,58,75,95,105,10aR,115,125)- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

ANSWER 18 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

DOCUMENT NUMBER: TITLE:

ANSVER 20 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
SSSION NUMBER: 2004:601035 HCAPLUS

MENT NUMBER: 142:169726

E: Site 1 sodium channel blockers prolong the duration of sciatic nerve blockade from tricyclic antidepressants

ORATE SOURCE: Department of Chemical Engineering, Massachusetts
Institute of Technology, Cambridge, MA, USA

Pain (2004), 110(1-2), 432-438

CODEN: PAINOBE ISSN: 0304-3959

ISHER: Besvier Ltd.

MENT TYPE: Journal AUTHOR(S): CORPORATE SOURCE:

SOURCE:

PUBLI SHER:

DOCUMENT TYPE: LANGUAGE: AB Many recei

JISHER: COURT PARKED 155M 0304-3959

ISHER: Blaevier Ltd.

MENT TYPE: Journal

MANDET TY

of drug injected at the sciatic nerve. In TCA-containing formulations, or blockade was consistently longer than thermal nociceptive block; motor blockade was also prolonged by tetrodotoxin and saxitoxin. In summary site 1 sodium channel blockers prolong the duration of TCAs via a locally mediated mechanism.

4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(site 1 sodium channel blockers tetrodotoxin combination with tricyclic antidepressants amitriptyline, nortriptyline and dowepin prolonged motor block was significantly longer than sensory block in rat)

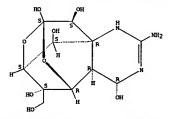
4368-28-9 HCAPLUS
5,917,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

ANSWER 20 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 21 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)



5

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 21 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:546375 HCAPLUS
111.99736 method and composition comprising local anesthetics and other agents for reducing reating membrane potential elec. disturbance, and use in organ preconditioning, arrest, protection, preservation and recovery.

INVENTOR(5): Dobson, Geoffrey Phillip Global Cardiac Solutions Pty Ltd, Australia PCCOMENT TYPE: Patent LANGUAGE: PIXXO2

DOCUMENT TYPE: Patent LANGUAGE: English DOCUMENT 11FE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English 2 PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004056181 Al 20040708 WO 2003-AU1711 20031222

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DW, DZ, EC, EE, EC, ES, ES, ET, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MY, MK, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SG, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TT, TT, TZ, UA, UG, US, UZ, VC, VN, VI, ZA, ZM, ZW, RW; BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CP, CG, CI, CM, GM, GM, GQ, GY, MM, MR, ES, SN, TD, TG
GB 2412067 Al 20050921 GB 2005-15048 20031222
US 2006034941 Al 20050921 GB 2005-15048 20031222
PRIORITY APPLN. INFO:

10 2006034941 Al 20050921 M2 2002-436175P P 20021223
AU 2003-903127 A 20030620
WO 2003-903127 A 20030620
WO 2003-903127 A 20030620
AD 2003-903127 A 20030620
anesthetic and of composition comprising an effective amount of a local anesthetic and of Composition comprising an effective amount of a local anesthetic and of Composition comprising an effective amount of a local anesthetic and of composition comprising an effective amount of a local anesthetic and other agents for reducing resting membrane potential velocity; USES (USes)

(Biological study); USES (USes)

(Biological study); USES (USes)

(CN 5,9:7,10a-Dimethano-loaH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-mino-1,4,4a,5,9,7,0-hexahydro-12-(hydroxymathyl)-,(4A,4aR,5B,75,55,5105,10aR,115,125)- (9CI) (CA INDEX NAME) APPLICATION NO. PATENT NO. KIND DATE DATE

Absolute stereochemistry.

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L8 ANSWER 22 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STM
ACCESSION NUMBER: 2004:546374 HCAPLUS
DOCUMENT NUMBER: 141:99735
TITLE: Compositions and methods using 1c
                                                                              141:99735
Compositions and methods using local anesthetics and other agents for organ preconditioning, arrest, protection, preservation and recovery
Dobson, Geoffrey Phillip
Global Cardiac Solutions Pty. Ltd., Australia
PCT Int. Appl., 150 pp.
CODEN: PIXXD2
Patent
English
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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PAT	ENT	NO.			KIN	O	DATE			APPL	ICAT				D	ATE		
wo	2004	0561	80		A1	-	2004	0708		WO 2		NU17			2	0031	222	
		AE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	œ,	CR,	cu,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	ĸж,	ΚZ,	ıc,	
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							ΗU,											
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PRIORITY	APP	LN.	INFO	.:					- 1	US 2	002-	(361	75P		P 20	0021	223	
										AU 2	003-9	9002	96		A 20	0030	123	
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AB The invention discloses a composition for arresting, protecting or preserving a cell, tissue or organ comprising an effective amount of a local anesthetic and of one or more of an anti-adrenergic, a calcium antagonist, an opioid, an NO donor and a sodium-hydrogen exchange inhibitor.

IT 4368-28-9, Tetrodotoxin
RI: PAC (Pharmacological activity), THU (Therapeutic use); BIOL (Biological study); USES (Uses) (compns. and methods using local anesthetics and other agents for organ preconditioning, arrest, protection, preservation and recovery)
RN 4368-28-9 HCAPLUS
CN 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

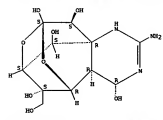
ANSWER 22 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

8

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN



REFERENCE COUNT:

THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 27

(Continued)

ANSWER 23 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

CAPIUS COFINGAL COCK.
2004:51404 ECAPUS
141:169243
Cardiovascular effects of the toxin(s) of the
Australian paralysis tick, Ixodes holocyclus, in the

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

PUBLISHER

LANGUAGE:

S):

Campbell, Fiona; Atwell, Rick; Fenning, Andrew; Hoey,
Andrew; Brown, Lindsay

TE SOURCE:

School of Veterinary Science, The University of
Queensland, Bribane, 4072, Australia

Toxicon (2004), 43(7), 743-750

CODEN: TOXIAG, ISSN: 0041-0101

ER:

Elsevier

TYPE:

Journal

E:

Extract of toxin(s) from the Australian paralysis tick, Ixodes
lus,

AB An extract of toxin(s) from the Australian paralysis tick, Ixodes holocyclus, produced pos. inotropic responses in rat left ventricular papillary muscles and pos. contractile responses in rat thoracic aortic rings. There was no measurable chronotropic response in rat right atria, but pos. inotropic conces. in papillary muscles produced arrhythmias in right atria. Fos. inotropic concess were attenuated by verapamil, but unaffected by metoprolol, cimetidine, pyrilamine, tetrodotoxin and pinacidil. Microelectrode studies on isolated left ventricular papillary muscles demonstrated that the extract prolonged action potential duration at 20, 50 and 90% of repolarization and delayed ventricular papillary muscle relaxation. Cardiovascular tissues isolated from rats with exptl. induced tick paralysis showed no myocardial damage as identified by histol. and ultrastructural examination The basal rate and force of contraction of isolated cardiact tissues were lower from tick-paralyzed than normal rats. Concentration-response curves to dobutamine and calcium chloride were

lar between tissues from tick-paralyzed and normal rats. Thus, the Australian paralysis tick, I. holocyclus, produces one or more towins with direct cardiovascular effects which mind the effects produced by direct blockade of cardiac and vascular K* channels.
4368-28-9, Tetrodotonin
RL: BSU (Biological study, unclassified), THU (Therapeutic use),
BIOL (Biological study), USES (Uses)
(paralysis tick toxins cardiovascular effects in rat and antiarrhythmic treatment)
4368-28-9 HCAPLUS

5,9:7,103-01methano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5,7,5,9,105,10aR,11S,128)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 24 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN 5510N NUMBER: 2004:486878 HCAPLUS 142:32376

ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

A novel toxicity-based assay for the identification of modulators of voltage-gated Na+ channels

AUTHOR(S): CORPORATE SOURCE: Weiser, Thomas

Weiser, Thomas
Boehringer Ingelheim Pharma GmbH & Co KG, Biberach,
D-88397, Germany
Journal of Neuroscience Methods (2004), 137(1), 79-85
CODEN: JNMEUT; ISSN: 0165-0270
Elsevier Science B.V. SOURCE:

PUBLI SHER:

CODEN: JMMEDT, ISSN: 0165-0270

ISHER: Blavvier Science B.V.
JOURNAL TYPE: Journal
JOURGE: English

Woltage-gated Na+ channels are promising drug targets. Screening of large
nos. of putative modulators, however, can be demanding and expensive. In
this study, a simpla, cheap, and robust assay to test the pharmacol.
modulation of Na+ channel function is presented. The assay makes use of
the fact that the intracellular accumulation of Na+ ions can be cytotoxic.
The toxicity of the Na+ channel activator veratridine in the presence of
an inhibitor of the Na+/KANTPase (ouabain) in a Navl.2a (cat brain IIA
e) expressing cell line is assessed. Na+ channel blockers should
reduce toxicity in this model. CHO cells which recombinantly expressed
rat Navl.2a subunits were seeded in 96-well plates, and cell survival was
tested after 24 h incubation in medium containing veratridine and ouabain in
the presence or absence of Na+ channel blockers. Propidium iodide
fluorescence was used as toxicity readout. Veratridine (100 µM) or
ouabain alone (500 µM) were not toxic to the cells. In the presence of
500 µM ouabain, however, veratridine induced half-maximal cell death
with an EC50 value of 15.1i2.3 µM. Ouabain's EC50 was 215.3i16.7
µM (vith 30 µM veratridine). The effects of a number of Na+ channel
blockers were tested and compared with their Na+ channel blocking activity
measured in voltage-clamp expts. Blockers from various chemical classes
reduced toxicity half maximally with IC50 values ranging from I.7a1.4

M (tetrodotoxin) to 280.5i48.0 µM (lamotrigine). There was a
linear relation between the log IC50 values obtained by the two methods
(slope: 1.180.08) correlation coefficient: 0.93). In summary, these data
show that this novel toxicity assay is well suited to test Na+ channel
blockers.
4368-2e-9, Tetrodotoxin

show that this novel toxicity assay is well suited to test Na+ channel blockers.

4368-28-9. Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Thexepeutic use); BIOL
(Biological study); USES (Uses)
(sodium channel inhibitor tetrodotoxin reduced toxicity half maximally and suppressed cell death in chinese hamster ovary cell transfected with rat brain type NaV1.2a subunit)
4368-28-9 BCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

L8 ANSWER 24 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 25 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

ANSWER 25 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
2004:486383 HCAPLUS
E: 2004:486383 HCAPLUS
Controlled-release pharmaceuticals for prolonged suppression of electrical activity in excitable tissues, and use in the treatment of epilepsy and other conditions
ANTOR(5): Kohane, Daniel S., Langer, Robert S.
Massachusetts Institute of Technology, USA; The General Hospital Corporation
CC: CC: PCT Int. Appl., 43 pp.
CODEN: PIXXO2
MEM TYPE: Patent
UMGE: Enelish

INVENTOR (5): PATENT ASSIGNEE (S):

SOURCE:

DOCUMENT TYPE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SR, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TC														TG				
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PRIORI'										US 21	002-	1302	10P	1	P 2	0021	202	
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activity in excitable tissues, and use in treatment of epilepsy and other conditions) 4368-28-9 HCAPLUS 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 26 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:475248 HCAPLUS DOCUMENT NUMBER: 141:238048

DOCUMENT NUMBER: TITLE:

AUTHOR (S): CORPORATE SOURCE:

141:238048
Prolonged infusion of tetrodotoxin does not block
mossy fiber aprouting in pilocarpine-treated rats
Buckmaster, Paul S.
Departments of Comparative Medicine and Neurology &
Neurological Sciences, Stanford University, Palo Alto,
CA. USA

CA, USA Epilepsia (2004), 45(5), 452-458 CODEN: EPILAK; ISSN: 0013-9580 Blackwell Publishing, Inc. SOURCE:

PUBLI SHER:

RCE: Epilepsia (2004), 45(5), 452-498

LISHER: CODEN: EPILEAK, ISSN: 0013-9580

LISHER: Blackwell Publishing, Inc.

UMENT TYPE: Journal

GUAGE: English

Mossy fiber sprouting is a common abnormality found in patients and models of temporal lobe epilepsy. The role of mossy fiber sprouting in epileptogenesis is unclear, and its blockade would be useful exptl. and perhaps therapeutically. Results from previous attempts to block mossy fiber sprouting and posttraums epileptogenesis. The present study tested the hypothesis that prolonged, focal infusion of tetrodotoxin would block mossy fiber sprouting and posttraums epileptogenesis. The present study tested the hypothesis that prolonged, focal infusion of tetrodotoxin would block mossy fiber sprouting after an epileptogenesit. The present study tested the hypothesis that prolonged, focal infusion of tetrodotoxin would block mossy fiber sprouting after an epileptogenic treatment. Adult rats were treated with pliocarpine treatment, a pump with a cannula directed toward the dentate gyrus was implanted to deliver 10 pM tetrodotoxin or vehicle alone at 0.25 µl/h. This method blocks local EEG activity in the hippocampus (Galvan et al. J Neurosci 2000; 20:2904-16). After 28 days of continuous infusion, rats were perfused with fixative, and their hippocampi analyzed anatomically with stereol. techniques. Tetrodotoxin infusion infusion, rats were perfused with fixative, and their hippocampi displayed similar levels of hilar neuron loss. The Timm stain revealed mossy fiber sprouting regardless of whether hippocampi were treated with tetrodotoxin infusion, vehicle infusion, or neither.

Frolonged infusion of tetrodotoxin dot block mossy fiber sprouting. This finding suggests that sodium channel-mediated neuronal activity is not necessary for mossy fiber sprouting after an epileptogenic treatment. 4368-28-9, Tetrodotoxin and prolonged infusion does not block mossy fiber sprouting in pilocarpine-treated rats)

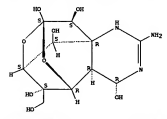
106-28-9-100-29-100-29-100-29-100-29-100-29-100-29-100-29-100-29-100-DOCUMENT TYPE: LANGUAGE: AB Mosey fibe

L8 ANSWER 26 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 27 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)



REFERENCE COUNT:

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 24

ANSWER 27 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN 2004:284577 HCAPLUS 40:368272

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

140:36272
Particular sensitivity to calcium channel blockers of the fast inward voltage-dependent sodium current involved in the invasive properties of a metastatic breast cancer cell line

Roger, Sebastien: Le Guennec, Jean-Yves: Besson, Pierre AUTHOR (5):

Pierre
Nutrition, Croissance et Cancer, Emi-U 0211, Faculte
de Medecine, Tours, 37032, Fr.
British Journal of Pharmacology (2004), 141(4),
610-615
CODEN: BJPCBM, ISSN: 0007-1188
Nature Publishing Group
Journal CORPORATE SOURCE:

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

NAME: Bnglish
A voltage-dependent sodium current has been described in the highly
invasive breast cancer cell line MDA-MB-231. Its activity is associated

Invasive breast cancer cell line NDA-MB-231. Its activity is associated the invasive properties of the cells. The aim of our study was to test whether this current (INa) is sensitive to three representative calcium channel blockers: verapami, ditiazem and nifedipine. INa was studied in patch-clamp conditions. INa was sensitive to verapamil (IC50 = 37.622.5 µM) and ditiazem (S3.221.6 µM), while it was weakly sensitive to nifedipine. The tetrodotoxin (TTX) concentration, which fully blocks INA (30 µM), did not affect cell proliferation. Ditiazem and verapamil, at concns. that do not fully block INA, strongly reduced cell proliferation, suggesting, regarding proliferation, that these mols. act on targets distinct from sodium channels. These targets are probably not other ionic channels, since the current measured at the end of a 500 ms long pulse in the voltage range between -60 and +40 mV was unaffected by verapamil and diltiazem. We conclude that the sodium channel supressed in MOA-MS-231 cells is sensitive to several calcium channel blockers. The present study also underlines the danger of concluding the possible involvement of membrane channel proteins in any phenomenon on the sole basis of pharmacol., and without an electrophysiol. confirmation.

4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THO (Thermacutic use); BIOL (Biological study); USES (Uses)

[sensitivity to calcium channel blockers of fast inward voltage-dependent yUSES (Uses)

[sensitivity to calcium channel blockers of fast inward voltage-dependent sodium current characteristic of metastatic breast cancer cells)

Cancer cells) 4168-28-9 HCAPUUS 5,9:7,10a-1; dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 28 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:151294 HCAPLUS DOCUMENT NUMBER: 140:368075 Synthesis of some name of some number of some name of some number of some name o

140:368075 Synthesis of some newer derivatives of substituted quinazolinonyl-2-oxo/thiobarbituric acid as potent

AUTHOR (5): CORPORATE SOURCE:

quinazolinony1-2-oxo/thiobarbituric acid as potent anticonvulsant agents
Archanar Srivastava, V. K.; Kumar, Ashok
Department of Pharmacology, Medicinal Chemistry
Division, L.L.R.M. Medical College, Meerut (U.P.),
250004, India
Bioorganic & Medicinal Chemistry (2004), 12(5),
1257-1264
CODEN: BMECEF; ISSN: 0968-0896
Elsevier Ltd.
Journal
English

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PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI English CASREACT 140:368075

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

5-{1'-[3''-Aminoacetyl-2''-methyl-6'',8''-dihalosubstitutedquinazolin-4''(3''H)-onyl]-thiosemicarbazido)-2-oxo/thiobarbituric acids and 5-{2'-amino-5'-{3''-aminoacethylene-2''-methyl-6'',8''-dihalosubstitutedquinazolin-4''(3''H)-onyl]-1',3',4'-thiadiazol-2'-yl)-2-oxo/thiobarbituric acid were prepared by incorporating 1-{3''-aminoacetyl-2''-methyl-6'',8''-dihalosubstituted-quinazolin-4'(3'H)-onyl)-thiosemicarbazides and 2-amino-5-{3''-aminoacetyl-2''-methyl-6'',8''-dihalosubstituted-quinazolin-4'(3'H)-onyl]-1,3,4-thiadiazoles resp. at 5th position of 2-oxo/thiobarbituric acids (via Mannich reaction). All the nevly synthesized compds. were screened for their anti-convulsant activity in MES and FTZ models and were compared with standard drugs phenytoin sodium and sodium valproate. Interestingly, these compds. were found to be devoid of sedative and hypnotic activities when tested. Out of the compds. studied, the most active compound I showed activity (90%) more potent than the standard drug. 683236-24-09
RI: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

RE: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(synthesis of quinazolinonyl-2-oxo/thiobarbituric acids as potent
anticonvulsant agents)
683236-24-0 HCAPLUS
Glycine, N-(2-methyl-4-oxo-3(4H)-quinazolinyl)-, 2-[[[(hexahydro-2,4,6trioxo-5-pyrimidinyl)methyl]amino]thioxomethyl]hydrazide (9CI) (CA INDEX

L8 ANSWER 28 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 29 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN PREF (Preparation); USES (Uses) (Continued)

(conjugates) hapten-carrier conjugates comprising a hormone, toxin, or drug and a core particle of bacteriophage protein for diagnosis and

the rapy) HCAPLUS

5.69-28-9 HCAPLUS

5.99-7, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4aR,5R,75,87,505,10aR,115,125)- [9CI] (CA INDEX NAME)

Pregant

L8 ANSWER 29 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:80526 HCAPLUS
DOCUMENT NUMBER: 140:144688
Hapten-carrier conjugates comprising hormone, toxin, or drug for diagnosis and therapy
Bachmann, Nartin F., Maurer, Patrik
Cytos Biotechnology Ag, Switz.
PCT Int. Appl., 144 pp.
CODEN: PIXXO2
DOCUMENT TYPE: Patent
English
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

		TENT						DATE										
	wo	2004	0091	16		A2		2004	0129		WO 2	003-	EP78	50		2	0030	718
	wo	2004	0091	16		A3		2004	0318									
		¥:	AE,	AG,	AL.	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	Cλ,	CH,	CN,
			co.	CR.	CU.	cz.	DE.	DK.	DH.	DZ.	EC.	EE.	ES.	FI.	GB.	GD.	GE.	GH,
									IS.									
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									SC,									
									UZ,							,		
		RW:							SD.							AM.	AZ.	BY.
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									GΑ,									
	CA	2487	849	,	,	AA	,	2004	0129	,	CA 2	003-	2487	849	,	2,	0030	718
	All	2487	2501	06		A1		2004	0200		AII 2	003-	2501	06		5	กกรก	718
	115	2004	0590	94		A1		2004	0325		115 2	003-	6220	64		5	กกรถ	718
	115	6932	971	•		B 2			0823		UJ 2		OLLU	••		~	0030	
		2003									RD 2	003-	1220	7		,	กกรก	718
	PD	1523	334	•		12		2005	0420		FD 2	003-	7650	47		5	กกรถ	718
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	ar.	2006																
	110	2005	2010	46		12		2000	1222		UF 2	005	1254	00		2	0050	F 1 0
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												003-	2070				$\alpha \alpha \alpha \alpha$	

The present invention provides compns. comprising a conjugate of a hapten with a carrier in an ordered and repetitive array, and methods of making such compns. The conjugates and compns of the invention may comprise a variety of haptens, including hormones, toxins and drugs, especially drugs

addiction such as nicotine. Compns. and conjugates of the invention are useful for inducing immune responses against haptens, which can use useful in a variety of therapeutic, prophylactic and diagnostic regimens. In certain embodiments, immune responses generated using the conjugates, compns. and methods of the present invention are useful to prevent or treat addiction to drugs of abuse and the resultant diseases associated with

treat addiction to drugs of abuse and the resultant diseases associated drug addiction.
4369-28-99, Tetrodotoxin
RL: ARU (Analytical role, unclassified); BPN (Biosynthetic preparation); BSU (Biological study, unclassified); DGN (Diagnostic use); TRU (Therapeuto use); ATT (Analytical study); BIOL (Biological study);

L8 ANSWER 30 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:80465 HCAPLUS
140:139471
TITLE: Preparation of of quinazolinone-like derivatives to

Preparation of of quinazolinone-like derivative: treat cellular proliferative diseases Bergnes, Gustave; Smith, Whitney W.; Yao, Bing; Morgans, David J., Jr.; MacDonald, Andrew Cytokinetics, Inc., USA PCT Int. Appl., 64 pp. CODEN: PIXXD2 Patent INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

		ENT						DATE			APPL	CAT	ION	NO.		D	ATE	
							-									-		
	WO	2004	0090	36		A2		2004	0129		WO 2	003~	US23	319		2	0030	723
	WO	2004	0090	36		A3		2004	0819									
		¥:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co.	CR.	CU.	CZ.	DE.	DK,	DM.	DZ.	EC.	EE.	ES.	FI.	GB.	GD.	GE.	GH.
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			UA,	UG,	υs,	UΖ,	vc,	VN,	YU,	ZA,	ZM							
		RW:	GH,	GM,	ΚE,	LS,	MW.	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	λZ,	BY,
			KG.	KZ.	MD.	RU.	TJ.	TM,	AT.	BE.	BG.	CH.	CY.	CZ.	DE.	DK.	EE.	ES.
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·/	EP	1537	089			A2		2005	0608	_	EP 2	003-	7660	28	•	2	0030	723
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	70	2006																
						12		2006	0112									
PRIO	JP 2006501201 RIORITY APPLN. INFO.:										US 2							
											WO 2	003-	US23	319	1	¥ 2	0030	723
OTHE	R SO	URCE	(S):			MAR	PAT	140:	1394	71								

R SOURCE(S): MARPAT 140:139471
The invention relates to quinazolinone-like derivs, that are inhibitors of the mitotic kinesin XSP and are useful in the treatment of cellular proliferative diseases, for example cancer, hyperplasias, restenosi; cardiac hypertrophy, immune disorders and inflammation. Preparation of 3-Benzyl-7-chloro-2-(3-benzyl-2-oxohexahydropyrimidin-4-yl)-3H-quinazolin-4-one is included.
651:232-36-3P
RI: PAC (Pharmacological activity): SPN (Synthetic preparation): TRU (Therapeutic use): BIOL (Biological study): PREP (Preparation): USES (Uses)

(Uses)
(preparation of quinazolinone derivs, to treat cellular proliferative diseases)
651923-36-3 HCAPLUS
4(3H)-Quinazolinone, 7-chloro-2-[hexahydro-2-oxo-3-(phenylmethyl)-4-pyrimidinyl]-3-(phenylmethyl)- (9CI) (CA INDEX NAME)

L8 ANSWER 30 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 31 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 31 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:62959 HCAPLUS
DOCUMENT NUMBER: 141:155568

An experimental vaccine against tetrodotoxin with longer term of validity

AUTHOR(S): Xu, Qinhuir Wei, Changhuar Huang, Kair Gao, Lishar Rong, Kangtair Yun, Liuhong
CORPORATE SOURCE: To divide Manyixue Zazhi (2003), 19(5), 339-342

CODEN: 2MZAER; ISSN: 1000-484X

PUBLISHER: CODEN: 2MZAER; ISSN: 1000-484X

PUBLISHER: Zhonguo Mianyixue Zazhi Bianjibu
DOCUMENT TYPE: Journal
LANGUAGE: Chinese

AB Objective: To develop an antitoxin vaccine against tetrodotoxin (TTX) and to explore the possibility of immune prevention and treatment for TTX intoxication. Methods: TTX was conjugated with Tachypleus tridentatus hemocyanin (TTH) in presence of formaldehyde and applied to immunize Balb/C mice. The level of antisers in the animals was periodically measured by ELISA and competition-inhibited enzyme immunoassay (CIBIA). Hice immunized with TTX-TTM were conjugated with emunoassay (CIBIA). Hice immunized with TTX-TTM were challenged ip. with low dones of TTX (1LD-13.5 Hg/kg). Results: The high titer and affinity of antisers lasted for as long as more than one year. The immunized mice were i.p. challenged with IXLD of TTX once and again at a fixed period, there was a affirmative antitoxic effect in about 12 mm (cotal ISALD), and a partial effect in following time. About one fourth of animal survived till 24 mo post initial immunization (cotal ZekuD), and which was a stage of sensecence in mice. The anti-TTX polsoning effect of animal was consistent with the antisers quality tested. Conclusions: The expl. vaccine of TTX could effectively protect animal from TTX intoxication and its effect was of longer duration of validity. Immunoprophylaxis would be the hopeful means for detoxification of TTX.

17 4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(Tachypleus tridentatus hemocyanin conjugates; tetrodotoxin vaccine with longer term of validity

Absolute stereochemistry.

L8 ANSWER 32 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
140:144682
Holecular antigen arrays comprising AP205 virus-like
particle and antigen for prevention and treatment of
cancer, drug addiction, poisoning, infection, and
allergy

allergy
Bachmann, Martin F.; Tissot, Alain; Pumpens, Paul;
Cielens, Indulis; Renhofa, Regina
Cytos Blotechnology AG, Switz.
PCT Int. Appl., 170 pp.
CODEN: PIXXD2
Patent INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA:	PATENT NO.						DATE								D	ATE	
															-		
WO	2004	0075	38		A2		2004	0122		WO 2	003-	EP75	72		2	0030	714
WO	2004	0075	38		A3		2004	0304							_		
	W:						AU,	A7.	RA.	RR.	BG.	RR.	RY.	R2	CA	CH	CN.
							DK,										
							IN,										
							MD,										
		PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	5G,	SK,	SL,	SY,	TJ,	TM,	TN,	
		TT,	TZ,	UA,	UG,	US,	UZ.	VC.	VN.	YU.	ZA.	ZM.	ZW				
	RW:	GH,													AM.	AZ.	RY.
							TM.										
							IE,										
		Br,					CH,										
	2489				AA		2004										
	2003						2004	0202		AU 2	003-	2466	90		2	3030 .	714
US	2004	0766	11		A1		2004	0422		US 2	003-	6178	76		2	0030	714
EP	1532	167			A2		2005	0525		EP 2	003-	7638	29		2	0030	714
		AT,															
							RO,										
RD	2003	0129	35	,			2005	0621	٠.,	DD 3	003-	1202		ш,	,	0020	774
PRIORITY					-		4003	0021									
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ORITY APPLM. IMFO:

US 2002-396126P P 20020717

Propresent invention provides a composition comprising an AP205 virus like particle (VLP) and an antigen. The invention also provides a process for producing an antigen or antigenic determinant bound to AP205 VLP. AP205 VLP bound to an antigen is useful in the production of compns. for inducing immune responses that are useful for the prevention or treatment of diseases, disorders or conditions including infectious diseases, allergies, cancer, drug addiction, potsoning and to efficiently induce self-specific immune responses, in particular antibody responses. Allergies, cancer, drug addiction, potsoning and to efficiently induce self-specific immune responses, in particular antibody responses. Allergies, Cancer, drug addiction, potsoning and to efficiently induce self-specific immune responses, in particular antibody responses.

Alle ABU (Analytical role, unclassified), BBN (Biosynthetic preparation);

BSU (Biological study, unclassified), DGN (Diagnostic use); TMD (Therapsutic use), NMST (Analytical study); BIOL (Biological study);

PREF (Preparation), USES (Uses)

(mol. antigen arrays comprising AP205 virus-like particle and antigen for prevention and treatment of cancer, drug addiction, poisoning, infection, and allergy)

4368-28-9 HCAPLUS

5,9:7,102-Dimethano-104H-[1,3]dioxocino[6,5-d)pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4s,5s,9,10-hexahydro-12-(hydroxymethyl)-,

(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9C1) (CA INDEX MAME)

L8 ANSWER 32 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

ANSWER 33 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Copentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4aR,5R,75,95,105,10aR,115,125)-(9CI) (CA INDEX NAME) (Continued)

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. PATENT NO. DATE OTHER SOURCE(s): MARPAT 140:99642 W9 2003-EF6665 V 20030625

The invention relates to novel medicament combinations based on sodium channel blockers and magnesium salts. The invention also relates to a method for the production thereof and the use thereof in the production of medicaments for the treatment of ischemic states. The sodium channel blockers and magnesium salts are administered parenteral) magnesium salts can be administered orally. The two components can be included in septormulations or in one formulation. Thus a sodium channel blocker injection contained (mg): crobenetine hydrochloride 7677 hydroxypropyl y-cyclodextrin 10000; mannicol 11000, acetic acid (99%) 152.557 sodium acetate trihydrate 56.55 and water to 250 mL. A magnesium salt injection contained 1000 mg magnesium sulfate and 10 mL water.

IT 4368-28-9, Tetrodotoxin
RL: TRU (Tharepeutic use); BIOL (Biological study); USES (Uses) (medicament combinations based on sodium channel blockers and magnesium salts)

ANSWER 34 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
SSION NUMBER: 2004:20436 HCAPLUS
E: Use of mixtures of related antigenic peptides to
induce a cytotoxic T lymphocyte immune response in a
wide range of individuals
NTOR(S): Ruprecht, Ruth M., Jiang, Shisong
Dana-Parber Cancer Institute, Inc., USA
PCT Int. Appl., 175 pp.
CCDEN: PIXXD2
MENT TYPE: Patent
UAGE: English INVENTOR (S) : PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004002415 A2 20040108 WO 2003-US20322 20030627

WO 2004002415 C2 20040603

WI AE, AG, AL, MM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GB, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MM, MH, MG, MK, MN, MM, MX, MZ, NI, NO, NZ, CM, PC, PH, PL, PT, RO, NG, US, UZ, VC, VN, YU, AZ, AM, ZW, KG, KZ, MD, RU, TJ, TH, AT, TZ, UA, UG, US, UZ, VC, VN, YU, AZ, AM, ZW, KG, KZ, MD, RU, TJ, TH, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CT, CM, CA, CM, ML, MR, NS, NT, DI TG

PRIORITY APPLN INFO: US 2004-22562 20041222

PRIORITY APPLN INFO: WO 2003-US20322 Al 20030627

AB The present invention provides compns. and methods for the treatment and prevention of immune disorders. A method of inducing an effective cytotoxic T lymphocyte (CTL) immune response in a wide range of individuals using mixts. of related antigenic pep ides (Overlapping Synthetic Peptide Formulations (OSFPS)) is described. OSFPs are derived from a longer antigenic peptide by splitting it up into peptides of at least eight amin acids with an overlap of at least one C-terminal amino acids with an overlap of at least one C-terminal amino acid from one peptide with the N-terminus of the next fragment. Use of an overlapping peptide library of the gag protein of HIV-I to induce CTL responses in BALB/c and GS7BL/6 mice is demonstrated. They also induced a proliferative T helper cell response.

IT 4868-28-9, Tetrodocsin

RL: ADV (Adverse effect, including toxicity), THU (Therapeutic uses), BIOL (Biological study); USES (USes)

(vaccines against, overlapping synthetic peptide formulations for; use of mixts. of related antigenic peptides to induce cytotoxic T lymphocyte immune response in wide range of individuals)

RN 4368-28-9 HCAPLUS

CN 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexaphy PATENT NO. KIND DATE APPLICATION NO. DATE

Absolute stereochemistry.

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

L8 ANSWER 34 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 35 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (4R, 4aR, 5R, 7S, 9S, 10S, 10aR, 11S, 12S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

HCAPLUS COPYRIGHT 2006 ACS on STN
2003:1006769 HCAPLUS
140:47530
Medicament combinations of sodium channel blockers and
fibrinolytics for treating ischemic conditions
Banzet, Sophie: Dusttmann, Hermann: Mauz, Annerose
Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.,
Germany
PCT Int. Appl., 29 pp.
CODEN: PIXXD2
Patent
German
: 1 L8 ANSWER 35 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: INVENTOR (S): PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE: LANGUAGE:

	PA	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION I	NO.		D	ATE	
							-									-		
	WO	2003	3105B	44		A1		2003	1224		WO 2	003-	EP58	13		2	0030	604
		¥:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co,	CR,	CU,	CZ.	DE.	DX.	DH.	DZ.	EC.	EE.	ES,	FI.	GB,	GD,	GE,	GH.
			GM,	HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	ĸ,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD.	MG,	MX,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ.	OH,
			PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	TJ,	TH,	TN.	TR.	TT,
			TZ,	UA,	UG,	US,	UZ,	vc.	VN,	YU,	ZA.	ZM,	ZW					
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD.	SL.	52.	TZ.	UG,	ZM,	ZW.	AM.	AZ.	BY,
								TM,										
								IE,										
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
	DE	1022	6814			A1		2004	0108		DE 2	002-	1022	6814		2	0020	615
	CA	2485	751			AA		2003	1224		CA 2	003-	2485	751		2	0030	604
		2003																
	EP	1515	720			A1		2005	0323		EP 2	003-	7599	07		2	0030	604
		R:	AT,	BE,	CH,	DE.	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	HC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
	JP	2005	5364	78		T2		2005	1202		JP 2	004-	5127	48		2	0030	604
	U5	2003	2355	76		A1		2003	1225	1	US 2	003-	1607	09		2	0030	612
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																	0020	
										1	70 2	003-1	EP58	13	1	j 2	0030	604

OTHER SOURCE(S):

ARRPAT 140:47530

AB The invention relates to novel medicament combinations based on sodium channel blockers and fibrinolytics, to a method for producing the same and to the use thereof for producing medicaments for treating ischemic conditions. The selected sodium channel blockers and fibrinolytics can be prepared as one formulation or as two formulations. The synthesis of benzazocine compds. that are sodium channel blockers is described. An injection formulation containing the sodium channel blocker included: crobenetine hydrochloride 767 mg; hydroxypropyl y-cyclodextrin 10000 mg; mannitol 11000 mg; acetic acid (99%) 125.25; sodium acetate trihydrate 56.6; water to 250 mL.

II 4368-28-9, Tetrodotoxin
RI: THU (Therspeutic use); BIOL (Biological study); USES (Uses) (medicament combinations of sodium channel blockers and fibrinolytics for treating ischemic conditions)
RN 4368-28-9 HCAPLUS

NN 4368-28-9 HCAPLUS

PR

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR (5):

ANSWER 36 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
25SION NUMBER: 2003:996337 HCAPLUS

LE: Analgesic effects of TTX alone and combined with morphine on formalin test in rats

NOR(S): Xu, Ying, Geng, Xingchao, Han, Jisheng, Qi, Shiquan, Xu, Baoshai

PORATE SOURCE: Department of Pharmacology, school of Basic Medical Sciences, Peking University, Beljing, 100083, Peop. Rep. China

NCE: Zhongguo Haiyang Yaowu (2003), 22(2), 39-41, 56

CODEN: ZHYARB; ISSN: 1002-3461

MENT TYPE: Shandongsheng Haiyang Yaowu Kexue Yanjiuso
Journal CORPORATE SOURCE:

SOURCE:

PUBLI SHER:

PUBLISHER: Shandongsheng Halyang Yaovu Kexue Yanjiuso DOCUMENT TYPE: Journal Journal LINGUAGE: Chinese AB The effects of tetrodotoxin (TTX) alone and combined with morphine on formalin-induced pain model were studied in rats. TTX, morphine, or both were administered i.m. and their effects were measured. Data were expressed as the median ID (IDSO). The IDSO of TTX alone was 0.8 µg kg-1. The IDSO of Morphine alone was 2.6 mg kg-1. The combination of TTX (39 µg kg-1 or 0.19 µg kg-1) and morphine showed more potent than each of them alone. The IDSO of Morphine reduced to 0.5 mg kg-1 and 1.1 mg kg-1, resp. An isobologram showed synergistic effect between TTX and morphine. The results indicated that TTX had analyssic effect in the formalin-induced pain model in rats. Comparing the effects of TTX alone and combined with morphine, the latter revealed a synergistic effect.

IT 458-28-9, Tetrodotoxin RI: PAC (Pharmacological activity), TMU (Therapeutic use), BIOL (Biological study), USES (Uses) (analyssic effects of TTX alone and combined with morphine on formalin test in rats).

test in rate)
4369-22-9 HCAPUM
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4a,8,7,5,9,105,10aR,118,125)- (9CI) (CA INDEX NAME)

L8 ANSWER 37 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:841693 HCAPLUS
141:135585
TITLE: 41:135585
Purification of tetrodotoxin with cationic exchange and gel filtration chromatograph for pharmaceutical and analytical use
INVENTOR(S): Jin, Chuanyin; Liu, Yongding; Song, Lirong; Zhu, Jimmine

PATENT ASSIGNEE(S):

Institute of Aquatic Biology, Chinese Academy of Sciences, Peop. Rep. China Faming Zhuanli Shenqing Gongkai Shuomingshu, 6 pp. CODEN: CROMEV SOURCE:

DOCUMENT TYPE: Patent

Chinese FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 1385432 A 20021218 CN 2001-114201 20010515

PRIORITY APPLM. INFO::

Method of the invention comprises adsorbing tetrodotoxin on the NH44-- or H++type weakly cationic exchange resin column, washing with water (buffer, or <0.4N accetic acid solution), eluting with 0.1-0.2N acetic acid as eluent or 0.01-0.1SN acetic acid as eluent, concentrating, dissolving in 0.01-0.1SN acetic acid (or picric acid); purifying on gel filtration column with 0.01-0.1SN acetic acid as eluent, and concentrating to obtain tetrodotoxin acetate (or tetrodotoxin picrate). The tetrodotoxin picrate may be converted into tetrodotoxin acetate by dissolving in water, precipitating

in NH4OH at pH 9, and dissolving in acetic acid.

RL: ARU (Analytical role, unclassified); BUU (Biological use, unclassified); PUR (Purification or recovery); TRU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses) (purification of tetrodotoxin with cationic exchange and gel filtration chromatograph for pharmaceutical and anal, use)

NS 9:17.10a-Dimethano-IoaH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 38 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
10:169452
TITLE:
Prolonged duration local anesthesis from tetrodotoxin-enhanced local anesthetic microspheres
Kohane, Daniel S.; Smith, Sarah E.; Louis, David N.;
Colombo, Gaias Ghoroghchian, Peter; Hunfeld, Nicole G.
M.; Berde, Charles B.; Langer, Robert
Hassachusetts Institute of Technology and Department of Anesthesis, and Research Associate, Massachusetts General Hospital and Harvard Medical School, Children's Hospital, Boston, MA, USA
Pain (2003), 104(1,2), 415-421
CODEN: PAINDB; ISSN: 0304-3959
Elsevier Science Ltd.
DOCUMENT TYPE:
JOURNAL BROWN TYPE:
AB There is interest in developing prolonged duration local anesthetics.
Here the authors examine whether tetrodotoxin (TTX) can be used to prolong the block from buptyscaine microspheres with and without dexamethasone.
Rats received sciatic nerve blocks with 75 mg of microspheres containing
O.051

0.05% (weight/weight) TTX, 50% (weight/weight) bupivacaine and/or 0.05% (weight/weight)

dexamethasone. 0.1% (weight/weight) TTX microspheres were also tested. The carrier fluid contained 1:100,000 epinephrine. Mociceptive and motor blockade of the hindpaw were quantified. Nerves and adjacent muscles were harvested 2 wk after injection for histol. assessment by light microscopy. The median nociceptive block duration in hours from the microsphere groups were: bupivacaine = 6.2, 0.05% TTX = 0. bupivacaine + TTX = 35.3, bupivacaine + dexamethasone = 31.3, TTX + dexamethasone = 8.1, TTX + bupivacaine + dexamethasone = 221.7. Some animals receiving particles containing 0.05% TTX had deficits in the uninjected extremity; all animals receiving 0.1% (weight/weight) TTX particles died. Pockets of particles were

receiving 0.1% (weight/weight) TTX particles died. Pockets of particles)
associated with localized inflammation, and all samples showed some evidence
of myotoxicity in the vicinity of the injection. The nerves themselves
appeared intact. In summary, coencapsulation of TTX in controlled release
devices containing bupivacaine and devamethasone resulted in very prolonged
nerve blocks. As formulated here, this preparation had a narrow margin of
safety. While the myotoxicity appears consistent with the well-known
reversible myotoxicity associated with local anesthetics, its long-term
significance remains to be established.

4368-28-9, Tetrodotoxin
RL: TMU (Therampeutic use): BIOL (Biological study): USES (Uses)
(prolonged duration local anesthesia from tetrodotoxin-enhanced local
anesthetic microspheres)

4368-28-9 HCAPUS
5,9:7,10a-Dimethano-10aH-[1,3]dicmocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(48,4ax,5x,75,95,105,10ax,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

(Continued) ANSWER 37 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 38 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 39 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:319348 HCAPLUS
DOCUMENT NUMBER: 138:331688
Methods of suppressing microglial activation and systemic inflammatory responses
Laskowitz, Daniel T.; Matthew, William D.; McMillian, McChael

PATENT ASSIGNEE(S): USA

PATENT ASSIGNEE(S): SOURCE:

USA
U.S. Pat. Appl. Publ., 48 pp., Cont.-in-part of U.S. Ser. No. 957,909.
CODEN: USXXCO

English 3

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 20030424 US 2003077641 US 2002164789 PRIORITY APPLN. INFO.: US 2002-252120 US 2001-957909 US 1998-77551P 20020923 20010921

US 1998-77551P P 19980311
US 1998-77551P P 19980311
US 1999-260430 B2 19990301
US 2001-957909 A2 20010921
Methods of suppressing the activation of microglial cells in the Central
Nervous System (CNS), methods of ameliorating or treating the neurol.
effects of cerebral ischemia or cerebral inflammation, and methods of
combating specific diseases that affect the CNS by administering a compound
that binds to microglial receptors and prevents or reduces microglial
activation are described. ApoE receptor binding peptides that may be used
in the methods of the invention are also described, as are methods of
using such peptides to treat peripheral inflammatory conditions such as
sepsis. Also described are methods of screening compds. for the shilty
to suppress or reduce microglial activation. Injection of ApoE (133-149)
in mice suppressed serum levels of TNFs and IL-6 following LPS
administration.
4369-28-9. Tetrodotoxin

in mice suppressed serum levels of TNFe and IL-6 following LPS administration.
4368-28-9, Tetrodotoxin.
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (BLOlogical study); USES (Uses) (anticonvulsant, ApoR receptor binding peptides suppressing microglial activation and systemic inflammatory responses)
4368-28-9 HCAPLUS
5,917, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,45,5,9,10-heraphylo-(12-(hydroxymathyl)-,(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 40 OF 108 HCAPFUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:196097 HCAPFUS
DOCUMENT NUMBER: 139:317174
TITLE: DOPA cyclohesyl ester potently inhibits
aglycemia-induced release of glutamate in rat striatal
slices AUTHOR (S) :

slices
Hashimoto, Mizukir Miyamae, Takeakir Yamamoto, Isao;
Goshima, Yoshio
Department of Molecular Pharmacology and Neurobiology,
Yokohama City University School of Medicine, Yokohama,
236-0004, Japan
Neuroscience Research (Oxford, United Kingdom) (2003),
45(3), 335-344
CODEM: NERADN; ISSN: 0168-0102
Elsevier Science Ltd.
Journal CORPORATE SOURCE:

SOURCE:

45(3), 335-344

CODEN: MERADN: ISSN: 0168-0102

PUBLISHER: Elsevier Science Ltd.
DOCUMENT TYPE: Journal

AB Brain inchemic insult causes glutamate release and resultant neuronal cell
death. We here show that L-3,4-dihydroxyphemylalanine (DOPA) is a pos.
regulatory factor for glutamate release elicited by a mild brain insult
using in vitro superfused rat striatal slices as a model system. Glucose
deprivation for 18 min elicited release of glutamate, DOPA and dopamine
(DA). Either tetrodotoxin (TTX) (1 mM) or a-methyl-p-tyroxine
(c-MPT) (1 mM), a tyroxine hydroxylase inhibitor reduced markedly
each of these releases. NSD-1015 (20 MM), an aromatic 1-amino acid
decarboxylase inhibitor restored the inhibition by c-MPT of
glutamate and DOPA but not DA release. DOPA cyclohexyl ester (DOPA CHE)
(0.3-1 MM), a competitive DOPA antagonist, concentration-dependently
suppressed aglycemia-induced glutamate release, the effect which was
minicked neither by S-sulpiride nor SCH23390, a DA DI or DZ receptor
antagonist, resp. Zonisamide (1-1000 MM), an anticonvulsant or TMB72
(1 MM), an e-amino-3-hydroxy-5-methyl-4-isomazole propionic acid
(AMPA) a receptor antagonist produced no effect on aglyceria-induced
glutamate release. DOPA CHE thus showed a relatively potent inhibitory
action on aglycenia-induced glutamate release among several
neuroprotective agents tested.

4569-28-9. Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(DOPA cyclohexyl ester potently inhibits aglycemia-induced release of
glutamate in rat striatum)
RN 4368-28-9. HCABLUS

N 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino(6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxysethyl)-,
(4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 39 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 40 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 39

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L8 ANSWER 41 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:76880 HCAPLUS
DOCUMENT NUMBER: 138:19602
TITLE: Methods of generating human cardiac cells and tissues
                                                  and uses thereof
Gepstein, Lior: Rehat, Izhak: Itskovitz-eldor, Joseph,
Amit, Michal
Technion Research and Development Foundation Ltd.,
INVENTOR(5):
PATENT ASSIGNEE(S):
                                                  Israel
PCT Int. Appl., 147 pp.
CODEN: PIXXD2
Patent
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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	PATENT NO.																
PAT	ENT I	ю.					DATE			APPL	ICAT	ION 1	NO.		_	ATE	
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	20030				A2		2003			WO 2	002-	I L60	6		2	0020	721
WO:	20030	0085	35		A3		2003	1023									
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	GM, HR, LS, LT,					MA.	MD.	MG.	MK.	MN.	NV.	MX.	MZ.	NO.	NZ.	OH.	PH.
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		KG,	ΚZ,	MD,	RU,	TJ,	TM,	AT,	BE.	BG.	CH.	CY,	CZ.	Œ,	DK.	EE.	ES.
		PI,	FR,	GB,	GR,	IE,	IT.	w,	MC,	NL,	PT,	SE,	SK,	TR,	BF,	BJ,	CF,
	FI, FR, CG, CI,						GQ.	GW.	ML.	MR,	NE.	SN,	TD.	TG			
US	20050						2005								2	0040	120
PRIORITY	APPI	N.	INFO	. :					- 1	US 2	001-	3064	52P		P 21	0010	720
									,	¥0 2	002-	I L60	5		A2 2	0020	721

A method of generating cells predominantly displaying at least one characteristic associated with a cardiac phenotype is displosed. The method comprises (a) partially dispersing a confluent cultured population of human stem cells, thereby generating a cell population including cell aggregates; (b) subjecting said cell aggregates to culturing conditions suitable for generating asheroid bodies; (c) subjecting said embryoid bodies; (c) subjecting said embryoid bodies to culturing conditions suitable for inducing cardiac lineage differentiation in at least a portion of the cells of said embryoid bodies, said culturing conditions suitable for inducing cardiac lineage differentiation including adherence of said embryoid bodies to a surface, and culture medium supplemented with serum, thereby generating cells predominantly displaying at least one characteristic associated with a cardiac phenotype.

predominantly displaying at least one characteristic associated with a cardiac phenotype.

4368-28-9, Tetrodotoxin
RL: PAC (Phareacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(methods of generating human cardiac cells and tissues and uses thereof)
4368-28-9 HCAPLUS
5,9:7, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4,5,9,10-hexabydro-12-(hydroxymethyl)-.
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

LO ANSWER	42 0	F 10	9 H	CAPL	US	COPY	RIGH	T 20	06 A	cs o	n ST	N				
ACCESSION N	UMBER	:		200	3:57	96	HCAP	LUS								
DOCUMENT NU	MBER:			138	: 499	52										
TITLE:				Use	of	iboe	um c	hann	el b	lock	ers	and	aspi	rin .	in	
				man	ufac	turi	na d	rugs	for	DEC	duci	ng a	nalo	esia		
									mamm			•				
INVENTOR(S)				Ku,	Bao	shan	, sh	um,	Hay	Kong						
PATENT ASSI	GNEE (5):		Wex	Med	ical	Ins	trum	enta	tion	Co.	, Lt	d.,	Peop	. Re	p.
				Chi	na											
SOURCE:				PCT	Int	. Ap	pl.,	11	pp.							
				COD	EN:	PIXX	D2									
DOCUMENT TY	PE:			Pat	ent											
LANGUAGE:				Chi	nese											
FAMILY ACC.			NT:	1												
PATENT INFO	RMATI	ON:														
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PATENT	NO.			KIN		DATE									ATE	
WO 200					_										0020	
WO 200				Cl		2003			# U 2	002-	CN42			2	0020	ora
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RW	: GH.								SZ.	TZ.	UG.	ZM.	ZV.	AM.	AZ.	BY
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	GN,	GQ,	GW,	ML,	MR,	NE,	5N,	TD,	TG							
CN 139				A		2003	0129		CN 2							
CA 249	3885			λA		2003	0103		CA 2	002-	2493	995		2	0020	618
ED 140						2004				~~~	7	~-		•		

CN 1393223 A 20030129 CN 2001-115990 20010622
CA 2493885 AA 20030103 CA 2002-2493885 20020618
EP 1405639 A1 20040407 EP 2002-754135 20020618
ER 147. BE, CH, UE, OK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CT, AL, TR
JP 2004534821 T2 2004118 JP 2003-506913 20020618
US 2004192659 A1 20040930 US 2004-480288 20040401
PRIORITY APPLN. INFO: CN 2001-115990 A 20010622
AB The present invention relates to the use of combinations of sodium channel blocking compds. and aspirin in manufacturing drugs for producing synergistically analgesic effect in mammals, in which said compds. bind to e-subunit of SSI or SS2 sites in the sodium channel. According to the invention, pharmaceutical compns. have enhancing analgesic effect, and therefore dosage of aspirin as well as its side effects would be reduced.

IT 4369-20-9, TTX
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(Use of sodium channel blockers and aspirin in manufacturing drugs for producing analgesia symergistically in mammals)

EN 4368-28-9 HCAPLUS
CN 5,917, Tola-Dimethano-10aH-[1,3]dicxocino(6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-besahydro-12-(hydroxymethyl)-,(4R,4am,5s,7s,9s,10s,10am,11s,12s)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 41 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 43 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 2002:905867 HCAPLUS HAPPENT NUMBER: 137:363099 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: 13/1303099 Analgesic composition and method Ku, Baoshan Shum, Frank Hay Kong Wex Medical Instrumentation Co., Ltd., Peop. Rep. INVENTOR (S) PATENT ASSIGNEE(S): China PCT Int. Appl., 35 pp. CODEN: PIXXD2 SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PA	ENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE		
							-									-			
	WO	2002	0942	72		A1		2002	1128		WO 2	002-	CN33	9		2	0020	520	
		v:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BΑ,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
			œ,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
			GH.	HR.	RU.	ID.	IL.	IN.	IS.	JP.	KE.	RG.	RP.	KR.	KZ.	LC.	LK.	LR.	
								HD,											
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								YU.										•	
		RV:						MZ.				TZ.	UG.	ZM.	ZW.	AT.	BZ.	CH.	
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	CN	1386						2002											
	US	2002	1982	26		A1		2002	1226	1	US 2	002~	6248	3		2	0020	205	
	us	6780	866			B2		2004	0824							_			
								2002			CA 2	002+	2485	337		2	0020	520	
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								ES,											
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	JP	2004						2004						89		2	0020	520	
								2004											
RIC												001-							
												002-							
												002-							

WO 2002-CN339 W 20020520
A pharmaceutical analgesic composition comprising an opioid analgesic agent and

A pharmaceutical analyssic composition comprising an optoid analyssic agent a compound that binds to the SSI or SS2 subunit of a sodium channel, such as tetrodotoxin and saxitoxin, and analogs thereof. Administration of an optoid analyssic agent and a compound that binds to the SSI or SS2 subunit of a sodium channel, such as tetrodotoxin and saxitoxin, and their analogs, produces analgesia in the treatment of pain in mammals. For example, the synergistic analgesia effect produced by co-administering tetrodotoxin (TTX) and morphine was observed in a formalin test in rats. Morphine used alone at 0.30 mg/kg only produced 10.2% inhibition of formalin-induced pain. Combination of TTX at 0.19 mg/kg with morphine at 2.50 mg/kg increased the inhibition rate to 86.7% from 34.9% where the latter was used alone. TTX at a dose of 0.39 mg/kg (1/50 of LDSO) produced an inhibition rate of 32.9% when used alone and 66.2% in combination with 0.15 mg/kg of morphine, whereas the latter only produced an inhibition rate of 7.2% when used alone. 4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Thermpeutic use); BIOL. (Biological study); USES (Uses)

L8 ANSWER 44 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:933513 HCAPLUS
DOCUMENT NUMBER: 137:304801
TITLE: Hethod of local anesthesia and analgesia using sodium channel blockers and local anesthetics
LLU, Yulingy Yin, Wenjuan
FATEAT ASSIGNEE(S): Wew Medical Instrumentation Co., Ltd., Hong Kong
U.S. Pat. Appl. Publ., 7 pp.
CODEN: USDNCC
DOCUMENT TYPE: Patent
LANGUAGE: Patent
EANGUAGE: 2nglish
FAMILY ACC. NUM. COUNT: 1

DATE APPLICATION NO. DATE

US 2002161013 A1 20021031 US 2001-6122 20011210
CN 1382443 A 20021204 (CN 2001-1010498 20010425
PRIORITY APPLM. INFO.: CN 2001-1101498 A 20010426
AB The invention relates to a method of obtaining local anesthesia and analyssis to the nerve tissue region of a mammal by administration of an ED of sodium channel blocking compds., including tetrodotoxin and/or saxitoxin and derivs. thereof, in a pharmaceutically suitable vehicle.

IT 4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therepeutic use); BIOL (Biological study); USES (Uses)
(method of local anesthesia and analgesia using sodium channel blockers and local anesthetics)
RN 4368-28-9 BTCAPLUS
CN 5,9:7,10a-0imethano-10aH-[1,3]dicasocino[6,5-d]pyrimidine-4,7,10,11,12-puntol, 2-mino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl).,
(48,4aB,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 43 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (synergistic analysis activity of combination of opioid and sodium channel blocker) 4368-28-9

\$100-20-7 nathus \$,9-7,10a-0imethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,87,87,5,95,103,10aR,115,128)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 45 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

2002:725238 HCAPLUS 138:281020

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

138:281020
Halothane attenuates the cerebroprotective action of several Na+ and Ca2+ channel blockers via reversal of their ion channel blockade
Oka, Michiko: Itoh, Yoshinori; Fujita, Takuya
Department of Biochemical Pharmacology, Kyoto
Pharmaceutical University, Kyoto, Yamashina, 607-8414, Janan

AUTHOR (S): CORPORATE SOURCE:

European Journal of Pharmacology (2002), 452(2), 175-181 SOURCE:

CODEN: EJPHAZ: ISSN: 0014-2999 Elsevier Science B.V.

PUBLISHER: DOCUMENT TYPE:

English

MENT TYPE: Journal NUGE: English The authors have previously shown the involvement of Na+ channel as well as N-type and P/O-type Ca2* channels in the oxygen and glucose deprivation-induced injury in rat cerebrocortical slices. In the present study, the authors investigated the influence of halothane on the cerebroprotective effects of a variety of Na+ and Ca2+ channel blockers in rat cerebrocortical slices. The hypoxic injury was attenuated by Na+ channel blockers including tetrodotoxin, lidocaine, and dibucaine, and Ca2+ channel blockers, such as verapamil, =-agatoxin IVA, and s-conotoxin GVIA. Halothane abolished the protective effects of lidocaine, dibucaine, and verapamil, all of which block the resp. cation channels in a voltage-dependent manner, without affecting the actions of tetrodotoxin, s-agatoxin IVA, and s-conotoxin GVIA, which reveal voltage-independent blockade. On the other hand, the NO synthesis estimated from the extracellular cyclic GMP formation was elevated during exposure to hypoxia. All channel blockers tested here attenuated hypoxia-evoked NO synthesis. Halothane blocked almost completely these actions of lidocaine and verapamil. No recover, the Na+ and Ca2+ channel blocked by these compds., as determined by veratridine- and XCI-stimulated

Synthesis, resp., was also reversed by halothane. These findings suggest that an anesthetic agent halothane reversed the Na+ and Ca2+ channel blockers of several voltage-dependent ion channel blockers, leading to the attenuation of their cerebroprotective actions. Therefore, the influence of halothane anesthesis should be taken into consideration for the evaluation of neuroprotective actions of Na+ and Ca2+ channel blockers.

4368-28-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USSS (Uses) (halothane on cerebroprotective effects of Na+ and Ca2+ channel blockers)

4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-loaH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol; 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R.4R,5R,75,95,105,10aR,115,125)- (SCI) (CA INDEX NAME)

L8 ANSWER 45 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

37 THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 46 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 46 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:408544 HCAPLUS
TITLE: 136:406875
THILE: 50406875
THILE: 50406875 LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2002041915 A1 20020530 WO 2001-CN1566 20011119
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DX, DM, DZ, EC, EE, SF, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LX, LR, LS, LT, LU, LV, MA, ND, MG, MK, NN, MW, MZ, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VW, YU, ZA, ZW, AM, AZ, BY, KG, KZ, LG, DW, TJ, TM, TR, CT, CD, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, ML, PT, SE, TR, BF, BJ, CP, CG, CI, CM, GA, GM, GO, GW, HL, MR, KE, SN, TD, TG
CN 1353990 A 20020619 CN 2000-132672 20001122
US 2002119987 A1 20020629 US 2001-181996 20010329
US 6559154 B2 20030506
AU 2002021491 A5 20020603 AU 2002-21491 20011119
EP 1335747 A1 20030820 BF 2001-997312 20011119
ER 3I, LT, LV, FI, RO, MK, CY, AL, TR
JP 2004513186 T2 20040503 JF 2002-344092 20011119
PRIORITY APPLN. INFO:
CN 2000-132672 A 20001122
AB The composition of the present invention comprises a sodium channel blocking compound which is capable of specifically binding to a site, either on an SS1 region or an SS2 region, on an extracellular region of a sodium channel alpha subunit, and a pharmaceutically acceptable carrier. An injection contained tetrodotoxin 1.5, 0.54 acetic acid 0.1, propylene glycol 80, and water for injection 100 mL. Stability of tetrodotoxin AL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmaceutical injections containing sodium channel blocking compds.)
RN 4368-28-9 H, CREDAUS APPLICATION NO. PATENT NO. KIND DATE

HAPPEN HCAPLUS CONSENTING CONTENTING CONTEN

Absolute stereochemistry.

L8 ANSWER 47 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
136:366124
Preparation of amidoalkyluracils as inhibitors of
poly(ADP-ribose)synthetase (PARS)
Albrecht, Barbara; Gerisch, Michael, Handke, Gabriele;
Jensen, Awel K Rahn, Thomas, Nickl, Werner; Oehne,
Felix; Schlemmer, Karl-Heinz; Steinhagen, Henning
Bayer Aq, Germany
CODEN: GWOKEX
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
CODEN: GWOKEX
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GERMAN FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	ENT	NO.					DATE				ICAT				D.	ATE		
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		1005															0001		
	CA	2428	335			AA		2002	0523		CA 2	001-	242R	335		2	0011	102	
	L.	2002	~~~					2002	0000										
											WO 2	001-	BP 12	694		2	0011	102	
	wo	2002	0404	55		C1		2002	0718										
		w:	AE.	AG.	AL.	AM.	AT.	AU.	AZ.	RA.	BB.	BG,	RR.	BY.	B7.	CA.	CH.	CNJ	
												EE,							
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP.	KE,	KG,	KP,	KR,	KZ.	LC.	LK.	LR.	
												MW,							
												SL,							
			υG,	US,	υz,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ.	MD,	RU.	ŤJ.	TM	
		RV:										TZ,							
												LU,						BF,	
			ВJ,	CF,	œ,	CI,	СМ,	Gλ,	GN,	GQ,	GW,	ML,	MR.	NE.	SN.	TD.	TG		
	UA	2002	0248	25		A5		2002	0527		All 2	002~	2482	5		2	0011	102	
	E.F	1339																	
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI.	LU.	NL.	SE.	MC.	PT.	
			TE.	SI.	LT.	LV.	FT.	RO.	MX.	CY.	AT.	TD							
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RIOR	UTI	APP	LN.	INFO	. :						DE 2	000-	1005	6312	1	۱ 2	3001	114	
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Iner	. 30	ONCE	(2):			MAK	LVI.	130:	2001	4									

Title compds. [1, λ = D, CH2D, DCH2, CH:CHCH2, CH2CH:CH, CH2CH2D, DCH2CH2, CH2DCH2; D = CH2, O, S; E, G = (substituted) alkylene, cycloalkylene; T =

ANSWER 47 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
CH2: U, V = (substituted) aryl, heterocyclyl; W = 0, 5, CO2, CCO, NR4; R4 = H, alkyl; m, n, q, p = 0, 1; X = 0, 5, NR5; R5 = H, alkyl; PhCH2; Y1 = H; Y2 = CH; Y1Y2 = 0, 5, NR6; R6 = H, alkyl; PhCH2; R1 = H, alkyl; PhCH2; PhC

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(preparation of amidoalkyluracils as inhibitors of poly(ADPribose) synthetase (PARS))
42653-30-9 MCAPLUS
1(2H)-Quinazolinepropanamide, 3,4,5,6,7,8-hexahydro-N-[2-[4-(7methylthieno[3,2-d]pyrimidin-4-yl)-1-piperazinyl]-2-oxoethyl]-2,4-dioxo(SCI) (CA INDEX NAME)

PAGE 1-A

L8 ANSWER 48 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STM
ACCESSION NUMBER:
DOCUMENT NUMBER:
136:325565
136:325565
Preparation of 3,4-dihydropyraindo(1,2-a)pyrimidines
and 3,4-dihydropyrazino(1,2-a)pyrimidines as

and 3,4-dinydropyrazino[1,2-ajpycimidines as analgesics gerlach, Matthias; Maul, Corinna; Jagusch, Utz-Peter Gruenenthal Gmbh, Germany PCT Int. Appl., 60 pp. CODEN: PIKKD2
Patent INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

								APPLICATION NO.										
WO									WO 2001-EP11702									
	₩:										, BG.							
											ES.							
											, KP,							
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											, w,							
											, ML,							
DE	1005										2000-							
AU	2002	0140	07		A5		2002	0422		AU	2002-	1400	7		7	0011	010	
ĊĀ	2425	685	•		AA		2003	0411		ca.	2002- 2001-	2425	685		5	0011	010	
EP	1325	010			A1		2003	0709		EP	2001-	9824	17		5	0011	010	
EP	1325	010			B1		2005	0427		_					_			
_										GR	, IT,	t.T.	IJI.	MT.	SE.	MC.	PT.	
	• • • •					PI.	RO.	MK.	CY.	AI.	TR							
BR	2001	0147	35		A.		2003	1014		RR	2001-	1473	5		2	0011	010	
JP	2004	5114	85		T2		2004	0415		JP	2002-	5343	20		2	0011		
NZ	5256	51			Ä		2004	1029		NZ	2002- 2001- 2001-	5256	51		2	0011		
AT	2941	80			E		2005	0515		AT	2001-	9824	17		2	0011		
ES	2239	168			т3		2005	0916		ES	2001-	1982	417		2	0011	010	
NO	2003	0015	88		Ä		2003	0408		NO	2001- 2003-	1588			2	0030	408	
US	2003	2203	22		Al		2003	1127		US	2003-	4096	14		2	0030	409	
ZA	2003	0036	34		A		2004	0812		2.8	2003- 2003-	3634			2	0030	512	
HK	1056	558			A1		2005	1216		HK	2003-	1089	15		2	0031	209	
PRIORIT	Y APP	LN.	INFO	. :						DE	2000-	1005	0661	- 1	N 2			
											2001-					0011		
OTHER S	OURCE	(S):			MAR	PAT	136:	32556		-								

OTH GI

AB Title compds. [I; Y = CR8; Z = N; or Y = N; Z = CR9; R1, R2 = H,

L8 ANSWER 47 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

PAGE 2-A

ANSWER 48 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (branched) (unsatd.) (substituted) alkyl, (unsatd.) (substituted) cycloalkyl, (unsatd.) (substituted) heterocyclyl, (substituted) (heterolaryl, (substituted) alkylaryl, etc. R3, R4 = H, H, (branched) (unsatd.) (substituted) alkylaryl, etc. R3, R4 = H, H, (branched) (unsatd.) (substituted) alkylaryl, etc. R3, R4 = Lc.; R5 = (branched) (unsatd.) (substituted) alkyl, (unsatd.) (substituted) cycloalkyl, (unsatd.) (substituted) heterocyclyl, (substituted) (heterolaryl, (substituted) heterocyclyl, (substituted) (heterolaryl, (substituted) alkylaryl, etc.; R6-R9 = H, F, Cl, Br, iodo, cyano, amino, aminoalkyl, aminodialkyl, etc.) and salts thereof vere prepd. Several I showed µ-opiate receptor binding with Ki = 1.4-2.5 µM and inhibited at 10 µM NDA/MX801 binding position with A0-478. The invention relates also to a method for the prodn. of the title compds., substance libraries conds. said compds., medicaments which contain said compds., the use of said compds. in the prodn. of medicaments for treating pain, urtnary incontinence, pruritus, tinnitus aurium and/or diarrhea and pharmaceutical prepns. contg. said compds.

RL: PAC (Pharmacological activity); SPM (Synthetic preparation); TBU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)
(Uses)
(preparation of dihydropyrimidopyrimidines and
dihydropyrazinopyrimidines as
analgesics)
RN 412350-23-3 HCAPLUS
CN 7,10-Nethano-31-pyrimido[1,2-a]quinazolin-3-one, 4,6,6a,7,10,10a-hexahydro1-hydroxy-2-methy1-6-(2-pyridiny1)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 49 OF 109 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:220374 HCAPLUS
DOCUMENT NUMBER: 136:241691
A method of analgesia using sodium channel blockers
Dong, Qingbin; Shum, Frank Haykong
WEM Medical Instrumentation Co., Peop. Rep. China
PCT Int. Appl., 60 pp.
CODEN: PIXXO2
PATENT TYPE: PRIXXO2 DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

P						KIN	D	DATE			APPI	ICAT	ION :	NO.		D.	ATE	
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¥	О	2002	0221	29		Al		2002	0321		WO 2	001-	CN13	91		2	0010	911
		w:	AE.	AG.	AL.	AH.	AT.	AU.	AZ.	BA.	BB.	BG,	BR.	BY.	BZ.	CA.	CH.	CN.
			co.	CR.	CU.	CZ.	DE.	DK.	DM.	DZ.	EC.	EE,	ES.	PT.	GB.	GD.	GE.	CH
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			02,	414	10,	٠.,	4w,	۸٦,	A2,	ы,	м,	KZ,	пD,	RU,	TJ,	TH		
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С	N	1356	104			Α		2002	0703		CN 2	000-	1245	17		2	2000	918
U	s	6407	088			B1		2002	0618		US 2	000-	6950:	53		21	J001(025
С	λ	2421	562			AΑ		2002	0321		CA 2	000- 000- 001- 002-	2421	562		21	0010	911
Α	U	2002	0137	85		A5		2002	0326		AU 2	002~	1378	5		21	0010	911
E	P	1320	369			A1		2003	0625		EP 2	001-	9820	91		21	0010	911
		R:	AT.	BE.	CH,	DE.	DK.	ES.	FR.	GB.	GR.	ĬΤ,	LI.	LU.	NL.	SE.	MC.	PT.
			TE.	SI.	LT.	LV.	FI.	RO.	MK.	CV.	AL.	ŤR						
В	R	2001	0139	61		A		2004	0113		BR 2	001-	1396	1		21	0010	911
J	P	2004	5084	04		T2		2004	0318		TP 2	002- 003-	5263	RO.		2	0010	911
E	Ŧ.	2003	0010	6		Ä		2005	0415		ER 2	003-	106			20	0010	911
	~	1563	839	-		A1		2005	0917	- 3	FD 2	004-	2202	9		- 2	0010	911
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	٠	2003	0003	12		•		2003	0423		NU 2	003-	312			21	7030	221
- 2	^	2003	0018	52		^		2004	0621		2A 2	003-	1825			20	2030.	306
N Z B RIORI	<u>ن</u>	10/6	90			٨		2004	0130		BG 2	003-	1076	90		20	1030	531
RIORI	ΤY	APP	LN.	INFO	.:					•	CN 2	000-	1245	17	1	1 20	30009	918
											EP 2	001-	98209	91	1	13 20	00109	911
										,	≌O 2	001-	CN 139	91	١	20	00109	911

This invention relates to a method of producing analgesia in a mammal experiencing pain by systemically administering an effective amount of a composition comprising essentially of a sodium channel blocking compound.

PR

suitable pharmaceutical vehicle, to alleviate the pain.
4366-28-9. Tetrodotomin
RL: ADV (Adverse effect, including toxicity), PAC (Pharmacological
activity), THU (Therapeutic use), BIOL (Biological study), USES
(Uses)
(analgesia using sodium channel blockers for neuropathic and cancer
nain)

pain) 4368-28-9 ECAPLUS 5,9:7,108-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-

L8 ANSWER 50 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:220373 HCAPLUS
171TLE: 1516:226908 A method of local anesthesia and analgesia using sodium channel blockers and local anesthetics
Ku, Baoshan; Qi, Shiquan
SOURCE: Ku, Baoshan; Qi, Shiquan
SOURCE: PATENT ASSIGNEE(S): 4CM Medical Instrumentation Co., Ltd., Peop. Rep. China
PCT Int. Appl., 25 pp. CODEN: PIXXID2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	ENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		0.	ATE	
						-									-		
WO	2002	0221	28		A1		2002	0321		WO 2	001-	CN13	90		2	0010	911
	W:	AE,	AG,	λĹ,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		α,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	15,	JP,	KE,	KG,	RP,	KR,	KZ,	LC.	LK,	LR,
		LS,	LT,	w,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,
		UZ,	٧N,	YU,	ZA,	ZW,	AM,	AΖ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TH		
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	5Ż,	TZ,	UG,	ZW,	AT,	BE,	Œ,	CY,
		ΟE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	w,	MC,	NL,	PŤ,	SE,	TR,	BF,
			CF,	Œ,	CI,	Οı,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG	
CN	1343	491			Α		2002	0410		CN 2	-000	1245	18		2	0000	918
US	6599	906			B1		2003	0729		US 2	-000	7028	26		2	0001	101
AU	2002	0137	94		A5		2002	0326		AU 2	002-	1378	4		2	0010	911
PRIORITY	APP	LN.	INFO	.:					1	CN 2	-000	1245	18		A 2	0000	918
									,	WO 2	001-	CN13	90	1	¥ 2	0010	911
AB The	pre	sent	inv	enti	on p	rovi	des	a me	thod	of	prod	ucin	g lo	cal .	anal	qesi	a an

The present invention provides a method of producing local analgesia and anesthesia in a mammal experiencing pain in a nerve tissue region. The method includes topically administering to the region, in a suitable pharmaceutical vehicle, an ED of a sodium channel blocking compound in a pharmaceutically suitable vehicle.

4368-28-9. Tetrodotomin
RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(local anesthesia and analgesia using sodium channel blockers and local anesthetics for neuropathic pain)
4368-28-9 ECAPLUS

5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(48,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX MAME)

Absolute stereochemistry.

ANSWER 49 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Copentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)-(9CI) (CA INDEX NAME) (Continued)

Absolute stereochemistry.

2

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 50 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 51 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

HCAPLUS COPYRIGHT 2006 ACS on STN
2002:72062 HCAPLUS
136:134774
Preparation of fused amidoalkyluracils as
poly(ADP-ribose) synthetase inhibitors
Haerter, Michaelr Albrecht, Barbaras Gerisch, Michaelr
Handke, Gabriele: Huetter, Joachims Jensen, Amelr
Krahn, Thomass Mittendorf, Joachims Oehme, Felixs
Schlemmer, Karl-Heinzs Steinhagen, Henning
Bayer Aktiengesellschaft, Germany
PCT Int. Appl., 113 pp.
CODEN: PIXX02
Patent INVENTOR (5) :

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent

PATENT NO. KIND DATE APPLICATION NO. DATE

US 6649618 PRIORITY APPLN. INFO.: DE 2000-10034801 WO 2001-EP7670

OTHER SOURCE(5): MARPAT 136:134774

Title compds. [I: A = D, CH2D, DCH2, CH:CHCH2, CH2CH:CH, CH2CH2D, DCH2CH2, CH2DCH2: D = CH2, O, S: X = (substituted) alkylene, cycloalkylene: R1 = H,

L8 ANSWER 52 OF 108 HCAPIUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:750642 HCAPIUS
DOCUMENT NUMBER: 135:284379

TITLE: P=scorpion toxin induces the release of y=iH_mainobutyric acid in rat brain slices
P=scorpion toxin induces the release of y=iH_mainobutyric acid in rat brain slices
P=scorpion toxin induces the release of y=iH_mainobutyric acid in rat brain slices
P=scorpion toxin induces the release of G=scorpion toxin induces the release of G=scorpion toxin induces the release of [3H]GABA from rat brain cortical slices is described. The stimulatory effect of TITX y on the release of [3H]GABA was dependent on incubation time and TITX y concentration with an ECSO of 0.19 µM. The scorpion toxin effect was Ca dependent and was completely inhibited by tetrodotoxin. P-Alanine also induced the release of [3H]GABA and this effect was not inhibited by tetrodotoxin but was additive in the presence of TITX y. The data suggest a neuronal origin for the release of (3H]GABA by TITX y.

If 4369-29-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study); USES (Uses)
(P=scorpion toxin induced release of y=sminobutyric acid in rat brain inhibition by tetrodotoxin)

N 4366-28-9 Tetrodotoxin
STAPIUS

N 5,9:7, 10a-10methano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-heashydro-12-(hydroxymethyl)-, (4R,4aR,5R,7s,9S,10S,10R,11S,12S)- (9CI) (CA INDEX NAME)

15

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 51 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (halogenated) alkyl, cycloalkylr R2 = SO2R4, SO2NR5R6, COR7, CONR8R9, COR7010; R4 = (substituted) alkyl, cycloalkyl, GE; E = (substituted) aryl, heterocyclyl, G is absent or (substituted) aryl, heterocyclyl, G is absent or (substituted) aryl, heteroarylr R5, R6 = H, (substituted) ycyloalkyl, aryl, heteroarylr GE (sa bore); R8, R9 = H, (substituted) alkyl, cycloalkyl, aryl or R1R2 = (substituted) heteroarylr R10 = (substituted) alkyl, cycloalkyl, aryl; or R1R2 = (substituted) monoor bicyclic heterocyclylr R3 = H, alkoxycarbonyl), were prepd. Thus, a mixt. of N-(3-aminopropyl)-N-benzyl-N-methylamine and tetrahydro-4H-thiopyran-4-one in PhMe was refluxed with camphorsulfonic acid followed by addn. of CiCONCO at room temp. to give 671 1-[3-benzyl(methyl) aminopropyl]-1,5,7,8-tetrahydro-2H-thiopyrano[4,3-d]pyrimidine-2,4(3H)-dione which was stirred with 2,2,2-trichloroethyl-Aloroformate in MeCN for 30 min at room temp. to give 631 2,2-trichloroethyl-3-[2,4-diono-3,4,7,8-tetrahydro-2H-thiopyrano[4,3-d]pyrimidin-1(SH)-yl)propyl(methyl)carbamate. Tested I showed 508 protection of endothelial cells with ECS = 0.05-0.5 µM.
390766-30-0P

RL: PAC (Pharmacological activity): SPN (Synthetic preparation): THU (Therapeutic use): BIOL (Biological study): PREP (Preparation): USES

(Uses)
(preparation of fused amidoalkyluracils as poly(ADP-ribose) synthetase inhibitors)
390766-30-0 HCAPLUS
2-Thiophenesulfonamide, N-[3-(3,4,5,6,7,8-hexahydro-2,4-dioxo-1(2H)-quinazolinyl)propyl]-N-methyl-5-[2-(methylthio)-4-pyrimidinyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 53 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN SSION NUMBER: 2001:748770 HCAPLUS HEAT NUMBER: 136:79620

DOCUMENT NUMBER: TITLE:

136:79620
Time course studies on the effectiveness of tetrodotoxin in reducing consequences of spinal cord

AUTHOR(5): CORPORATE SOURCE:

contusion
Rosenberg, Lisa J., Vrathall, Jean R.
Department of Neuroscience, Georgetown University,
Washington, DC, 20007, USA
Journal of Neuroscience Research (2001), 66(2),
191-202

SOURCE:

191-202 CODEN: JNREDK: ISSN: 0360-4012 Wiley-Liss, Inc.

PUBLI SHER:

PUBLISHER: Wiley-Liss, Inc.
JOURNAT TYPE: Journal
ANGUAGE: English
AB Focal injection of the sodium channel blocker tetrodotoxin (TTX) into the
injury site at either 5 or 15 min after a standardized thoracic contusion
spinal cord injury (SCI) reduces white matter pathol. and loss of axons in
the first 24 h after injury. Focal injection of TTX at 15 min after SCI
also reduces chronic white matter loss and hindlimb functional deficits.
We have now tested the hypothesis that the reduction in chronic deficits
with

TTX treatment is associated with long-term preservation of axons after SCI and compared both acute (24 h) and chronic (6 wk) effects of TTX administered at 15 min prior to and 5 min or 4 h after SCI. Our results indicate a significant reduction of acute white matter pathol. in rats

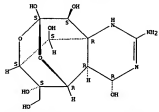
administered at 15 min prior to amp a min or 1 metros. Act treated
indicate a significant reduction of acute white matter pathol. in rats
treated
with TTX at 15 min before and 5 min after injury but no effect when
treatment was delayed until 4 h after contusion. Compared with injury
controls, groups treated with TTX at 5 min and 4 h after injury did not
show a significant deficit reduction, nor was there a significant sparing of
white matter at 6 wk compared with injury controls. In contrast, the
group treated with TTX at 15 min before SCI demonstrated significantly
reduced hindlimb functional deficits beginning at 1 wk after injury and
throughout the 6 wk of the study. This was associated with a significantly
higher axon d. in the ventromedial white matter at 6 wk. The results
demonstrate that blockade of sodium channels preserves axons from loss
after SCI and points to the importance of time of administration of such
drugs for therapeutic effectiveness.
IT 4360-20-9, Tetrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USBS (Uses)
(time course studies on effectiveness of tetrodotoxin in reducing
consequences of spinal cord contusion)
RN 4360-20-9 RCAPIUS
CN 5,9:7,10a-pimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-mino-1,4.4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7s,9s,10s,10aR,111s,125)- (SCI) (CA INDEX NAME)

L8 ANSWER 53 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 54 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)



REFERENCE COUNT:

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 54 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:660029 HCAPLUS DOCUMENT NUMBER: 136:335125

DOCUMENT NUMBER: TITLE:

High concentrations of adrenergic antagonists prolong sciatic nerve blockade by tetrodotoxin Kohane, D. S.; Lu, N. T.; Crosa, G. A.; Kuang, Y.; AUTHOR(S):

CORPORATE SOURCE:

Department of Anesthesia, Children's Hospital, Boston, MA, USA SOURCE: Acta Anaesthesiologica Scandinavica (2001), 45(7), 899-905

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

Acta Anaesthesiologica Scandinavica (2001), 45(7), 899-905
CODEM: AANEAB, ISSN: 0001-5172
Munker TYPE: Journal Munker International Publishers Ltd.
JOURNAL TYPE: Journal State of the State of State of

appear to be adrenergic-receptor-specific, or mediated by GTPase activation.
4368-28-9. Terrodotoxin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(adrenergic antagonists prolong sciatic nerve blockade by tetrodotoxin)
4368-28-9 HCAPLUS

4368-28-9 HCAPLUS 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 55 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:566880 HCAPLUS
DOCUMENT NUMBER: 135:134288
TITLE: Diagnostic kit and method and creatine recognizing
agents for detecting creatine levels
Al Athel, Fahad Mohammed Saleh Bell, Thomas W.,
Khasanov, Alisher B., Kaddurah-Daouk, Rima
Fal Diagnostics, USA
SOURCE: Patent
LNNGUAGE: PIXCOZ
DOCUMENT TYPE: Patent
English

English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA:	ENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D.	ATE		
						-									-			
WO	200	10557	19		A2		2001	0802		WO 2	001-	US26	50		2	0010	126	
WO		10557					2001											
	٧:	ΑE,	λG,	λL,	λM,	λT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH.	GM,	HR.	
		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	ıc,	LK.	LR,	LS.	LT.	
							MK,											
		SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TH,	TR,	ΤŤ,	TZ,	UA,	UG,	UZ,	VN,	YU,	
		ZA,	Ζ¥,	AM,	λZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TH						
	RW:	: GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	Z₩,	AT,	BE,	CH,	CY,	
		DE,	DX,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	
		BJ,	CF,	œ,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
บร	6566	6086			B1		2003	0520		US 2	000-	4942	05		21	0000	128	
ORIT	API	PLN.	INFO	.:						US 2	000-	1942	05	- 1	A 21	0000	128	
ER S	DURCE	E(S):			MAR	PAT	135:	1342	8									

Methods for the detection of creatine compound levels in body fluid samples are discussed. Portable kits capable of determining creatine levels using non-invasive and visually detectable methods are also included. I was prepared from quinaldine and used to detect creatine by IH NMR spectroscopy. 352229-25-5

RL: ARG (Analytical reagent use); THV (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (diagnostic kit and method and creatine recognizing agents for detecting creatine levels)

352229-25-5 HCAPLUS
[1,10]Phenanthrolino[2,3-b][1,10]phenanthroline-2-carboxylic acid, 13-{5-(1,3-4,6,11,11-hexahydro-2H-pyrindio(2,1-b)quinazolin-10-yl)ethynyl]-2-pyridinyl]-5,6,9,10-tetrahydro- (9CI) (CA INDEX NAME)

L8 ANSWER 55 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 56 OF 108 HCAPIUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 14

LS ANSVER 56 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:563098 HCAPLUS
DOCUMENT NUMBER: 137:122

Author (1):

Author (2):

Author (3):

CORPORATE SOURCE: Department of Blomonitoring and Sensoring, University Centre for Pharmacy, University of Groningen,
Groningen, 9713 AV, Neth.

SOURCE: JOURNAL OF SENSORIES English
AB For interpretation of microdialysis expts. in which compds. are applied by retrodialysis, it is important to have information about the migration rate of the infused compds. Here we describe a dual-probe microdialysis method that can be used to evaluate the penetration about the migration rate of the infused compds. Here we describe a dual-probe microdialysis method that can be used to evaluate the penetration rate of the infused distance (1 mm) of the infusion probe. Using this approach several compds., each known to induce specific changes in the extracellular levels of dopamine, were infused into the striatum of the rat. The results indicate that the penetration rate of the pharmacol. effect of recorded by a second probe positioned at a fixed compds., each known to induce specific changes in the extracellular levels of dopamine, were infused into the striatum of the rat. The results indicate that the penetration rate of the pharmacol. effect of infused compds. differed widely. No effects were seen at the second probe when high potassium chloride was infused. Apparently dopamine was not able to migrate into brain tissue over a distance of 1 mm. Low penetration rates were observed for the dopamine antagonist sulpride, the dopamine against LY 171555, and for amphetamine and nomifensine. A very high penetration rate was observed in case of tetrodotoxin (TTX). The fast effects of TTX could also be explained by resort inhibition of neurons passing along the infusion probe. The present study showed that most of the compds. have rather slow infusion rates, indicating that relatively high infusion concess. are needed (1-10 mm) to reach substantial brain concess. at a distance of 1

Absolute stereochemistry.

ANSWER 57 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SION NUMBER: 2001:453494 HCAPLUS 135:41024

DOCUMENT NUMBER:

TITLE:

.>3:4UL4 Compositions, kits, apparatus, and methods for inhibiting cephalic inflammation by intranasal administration of long-acting local anesthetic Levin, Bruce H.

INVENTOR(S): PATENT ASSIGNEE (S):

USA
U.S. Pat. Appl. Publ., 48 pp., Cont.-in-part of U.S.
Sec. No. 118,615.
CODEN: USXXCO SOURCE:

DOCUMENT TYPE: English 6

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	AP	PLICATION NO	·.	DATE

US 2001004644	A1	20010621	บร	2000-737302		20001215
US 2001055607	A1	20011227	US	1998-118615		19980717
US 6432986	B2	20020813				
US 2002010194	A1	20020124	US	2001-775724		20010201
US 2003133877	A1	20030717		2002-218138		20020812
US 2005281751	A1	20051222		2005-126475		20050511
PRIORITY APPLN. INFO .:			US	1997-90110P	P	19970721
				1998-72845P		19980128
				1998-84559P		19980506
				1998-118615		19980717
				1999-170817		19991215
				1997-897192		19970721
				1999-117398		19990127
				2000-492946		20000127
			US	2000-737302	B2	20001215
			US	2002-218138	A2	20020812
AB Methods, kits, app	paratus.	and compns.	for	inhibiting	cephalic	inflammati

Methods, kits, apparatus, and compns. for inhibiting cephalic inflammation, including meningeal inflammation and cerebral inflammation for example, in a human patient are provided. The methods comprise intranasally administering to the patient a pharenaceutical composition comprising a local anesthetic, and preferably a long-acting local anesthetic ingredient. A composition useful for practicing the methods of the invention is described which comprises at least one local anesthetic in a pharmaceutically acceptable carrier, wherein the composition is formulated for intranasal delivery. A kit comprising the composition and an intranasal applicator is also included in the invention. Apparatus for delivering or applying the composition of the invention or for performing the methods of the invention are also described. Ropivacaine was dorsonasally administered to individual patients experiencing head pain, other symptoms, or both, believed to be associated with an acute nigraine episode. Dorsonasally administered ropivacaine rapidly inhibited of migraine in 921 of the ambulatory patients, as evidenced by an average 901 reduction in perceived

within one hour, usually within 15 min or less. Symptoms of nausea and photophobia associated with acute migraine episodes in patients were similarly inhibited. Rebound of migraine occurred in only 5.4% of patients within twenty-four hours of treatment.

4368-28-9. Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse): BSU (Biological study, unclassified): THU (Therapeutic use): BIOL (Biological study): USES (Uses)
[compns. and kits and apparatus and methods for inhibiting cephalic inflammation by intransal administration of long-acting local

ANSWER 57 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

anesthatic)
438-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-heashydro-12-(hydroxymathyl)-,
(4R,4aR,S,7,9,5,105,10aR,115,125)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 58 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (4R,4aR,5R,75,95,105,10aR,115,125) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 58 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
SSION NUMBER: 2001:452849 HCAPLUS
HENT NUMBER: 135:56081
E: Compositions, kits, apparatus, and methods for inhibiting cephalic inflammation resulting in acute migraine and other paintl episodes associated with neurowascular disorders ACCESSION NUMBER: DOCUMENT NUMBER: INVENTOR (S): PATENT ASSIGNEE (S): SOURCE: Levin, Bruce H. USA PCT Int. Appl., 119 pp. CODEN: PIXXD2 Patent DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. PATENT NO. KIND DATE PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2001043733 A2 20010621 WO 2000-US33916 20001215

WO 2001043733 A3 20020510

W: AR, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CM, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GH, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, KK, M, MY, MX, MZ, MD, MZ, FL, PT, RD, RU, 2A, ZW

RW GH, GH, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GB, IE, IT, LU, NC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GM, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO: US 1999-170817P P 19991215

M Methods, kits, apparatus, and compns. for inhibiting cephalic inflammation, including meningeal inflammation and cerebral inflammation for example, in a human patient are provided. The treatment is for acute cerebral neurovascular disorders resulting in acute migraine and other painful episodes. The methods comprise intranasally administering to the patient a phareaceutical composition comprise intranasally administering to the patient a phareacturical composition comprise at least one local anesthetic in a pharmaceutically acceptable carrier, wherein the composition if formulated for intranasal delivery. A kit comprising the composition and an intranasal applicator is also included in the invention. Apparatus DATE and an intranasal applicator is also included in the invention. Apparatus delivering or applying the compns. of the invention or for performing the methods of the invention are also described.
4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TRU (Therapeutic use); BIOL (Biological study); USES (Uses)
study); USES (Uses)
[Compns., kits, apparatus, and methods for inhibiting cephalic ammation.

ammation
resulting in acute migraine and other painful episodes associated with
neurovascular disorders)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethane-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl)-,

ANSWER 59 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

SSION NUMBER: 2000:454869 HCAPLUS

131:222658
E: Inhibition of different pathways influencing Na+
homeostasis protects organotypic hippocampal slice
cultures from hypoxic/hypoglycemic injury

OR(S): Breder, J., Sabelhaus, C. F., Opitz, T., Reymann, K.
G.; Schroder, U. H.

ORATE SOURCE: Neurobiology, Magdeburg, D-39008, Germany
Neurobiology, Magdeburg, D-39008, Germany

CE: Neurobaranacology (2000), 39(10), 1779-1787

CODEN: NEPHBW; ISSN: 0028-3908 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: AUTHOR (S): CORPORATE SOURCE: SOURCE: PUBLI SHER: Elsevier Science Ltd.

LISHER: Elsevier Science Ltd.

MRNM TYPE: Journal

BUAGE: English

A prominent feature of cerebral ischemia is the excessive intracellular
accumulation of both Na+ and Ca2+, which results in subsequent cell death.
A large number of studies have focused on pathways involved in the increase
of the intracellular Ca2+ concentration (Ca2+); whereas the elevation of
intracellular Na+ has received less attention. In the present study we
investigated the effects of inhibitors of different Na+ channels and of
intracellular Na+ has received less attention. In the present study we
investigated the effects of inhibitors of different Na+ channels and of
ischemic damage in organotypic hippocampal slice cultures. The
synaptically worked population spike in the Ca1 region was taken as a
functional measure of neuronal integrity. Neuronal cell death was
assessed by propidium iodide staining. The Na+ channel blocker
tetrodoxoin, and the NMDA receiptor blocker MK 801, but not the
AMPA/kainate receptor blocker MRSO prevented ischemic cell death. The
novel Na+/Ca2+ exchange inhibitor 2-{2-{4-(4-introbenzyloxy) phenyl}ethyl]i
sothiourea methanesulfonate (KB-R7943), which preferentially acts on the
reverse mode of the exchanger, leading to Ca2+ accumulation, also reduced
neuronal damage. At higher concns. KB-R7943 also inhibits Ca2+ extrusion
by the forward mode of the exchanger and exaggerates neuronal cell death.
Neuroprotection by KB-R7943 may be due to reducing the [Ca2+)1 increase
caused by the exchanger.
4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study), USES (Uses)

(inhibition of different pathways influencing Na+ homeostasis protects
organotypic hippocampal slice cultures from hypoxic/hypoglycemic
injury)

4368-28-9 HCAPLUS

inflammation

organosypte inpposaspas 31cc collects ...,pas...,pas...,y,33..., injury)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl)-,
(48,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

L8 ANSWER 59 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 60 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

HCAPLUS COPYRIGHT 2006 ACS on STN 2000:351162 HCAPLUS 133:790 ANSWER 60 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: New use of glutamate ar cancer Ikonomidou, Hrissanthi use of glutamate antagonists for the treatment of INVENTOR(S): PATENT ASSIGNEE(S): Ikonomidou, Hrissanthi Germany Eur. Pat. Appl., 21 pp. CODEN: EPXXDW Patent English 1 DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. APPLICATION NO. KIND DATE DATE ## PAIL CATION NO. DATE | PAPEL CATION NO. PAPEL CATION NO

19931022

IE, FT, CY
US 6797692
B1 20040928
US 2001-830354
US 2005054619
A1 20050310
US 2004-912159
20040806
US 2005054650
A1 20050310
US 2004-912159
20040806
US 2005054650
A1 20050310
US 2004-91215
20040806
CRITY APPLN. INFO.:

EP 1998-250380
A 19981028
EP 1999-52622
A3 19991022
W0 1999-EP8004
V 1999-D2804
V 1999-D2804
A3 20010425
New therapies can be devised based upon a demonstration of the role of glutamate in the pathogenesis of cancer. Inhibitors of the interaction of glutamate with the AMPA, kainate, or NMDA receptor complexes are likely to be useful in treating cancer and can be formulated as pharmaceutical compns. They can be identified by appropriate acreens.
4368-28-9, Tetrodotowin
RE: BAC (Biological activity or effector, except adverse); BSU (Biological study), unclassified); TMU (Therapeutic use); BIOL (Biological study); USES (USES)
(glutamate antagonists for cancer treatment)
4368-29-9 HCAPLUS
5,917,10a-Dimethano-10aH-[1,3]dioxocino[6.5-4]

4368-28-9 HCAPUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 61 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: DOCUMENT NUMBER: 2000:290862 HCAPLUS 132:303512 132:303512
Methods for enhancing wound healing
Gassner, Holger G.; Sherris, David A.
Mayo Foundation for Medical Education and Research, TITLE: INVENTOR (5): PATENT ASSIGNEE (S): USA PCT Int. Appl., 24 pp. CODEN: PIXXD2 SOURCE: DOCUMENT TYPE: Patent LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English WO 2000024419 A1 20000504 WO 1999-US24182 19991015

V: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JF, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MM, MX, NO, NZ, PL, PT, NO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, AZ, ZW

RY: GH, GM, KE, LS, MY, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GM, GW, ML, HR, NE, SN, TD, TG

CA 2347828 A2 20000504 CA 1999-2347828 19991015

EP 1128844 A1 20010505 EP 1999-960130 19991015

EP 1128844 A1 20010505 EP 1999-960130 19991015

EP 1128844 B1 20060104

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, CY

JF 2002528421 T2 20020910 US 2001-897072 19991015

US 2003036502 A1 20020910 US 2001-897072 20010418

US 2005175637 A1 20020911 US 2005-61299 20050218

PRIORITY APPLN. INFO::

US 1998-105688P P 19991027 US 2006039930 A2 2006023

RITY APPLN. INFO::

US 1998-105688P P 19981027

W0 1999-US24182 W 19991015

US 2001-89702 A1 20011126

A method for treating a patient having a wound is described. The method includes administering an amount of a chemodenervating agent such that healing of the wound is enhanced. The method is illustrated by detailing the mean differences of the scores of the paired exptl. and control scars across three observers. Also claimed is a local administration of compns. containing chemodenervating agents (e.g. botulinum toxins), local thetics

containing chemodenervating agents (e.g. potulinum coming), local thetics (e.g. lidocaine), and vasoconstrictors (e.g. epinephrine). 4369-28-9, Tetrodotoxin RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THO (Therapeutic use); BIOL (Biological study); USES (Uses) (local administration of compns. containing chemodenervating agents and anesthetics and vasoconstrictors for enhancing wound healing) 4368-28-9 HCAPLUS 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,SR,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

ANSWER 61 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERÊNCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 62 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (4R, 4aR, 5R, 7s, 9s, 10s, 10aR, 11s, 12s) - (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 62 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2000:229542 HCAPLUS

132:288238 DOCUMENT NUMBER: TITLE:

Z0001:22932 RAPLOS
Organotypic hippocampal slice cultures as an in vitro
model for the investigation of neuroprotective drugs
against ischemic damage
Breder, Jorg Sabelhaus, Clemens F., Schroder, Ulrich
H., Reymann, Klaus G.
Laboratory of Neuropharmacology, Leibniz Institute for
Neurobiology, Magdeburg, 39008, Germany
Schriften des Forschungszentrums Juelich,
Lebenavissenschaften/Life Sciences (1999), 3(Cell
Culture Models as Alternatives to Animal
Experimentation for the Testing of Neuroprotective
Compounds in Stroke Research), 79-98
CODEN: SFLSF9, ISSN: 1433-5549
Forschungszentrum Juelich GmbH
Journal
English

CORPORATE SOURCE:

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

AUTHOR(S):

MENT TYPE: Journal
SUAGE: Forschungszentrum Juelich GmbH
JUGE: English
Cerebral ischemia results in severe cell degeneration and consequently in
loss of brain functions. In animal models of global ischemia the
hippocampus has turned out to be 1 of the most vulnerable brain areas, and
within the hippocampus the pyramidal neurons of the CAI region are highly
susceptible. These in vivo test systems cause substantial stress in form
of pain and anxiety to the animals involved, giving rise to ethical
problems and little public acceptance. In vitro models were developed to
overcome these problems. Dissociated cell cultures allow the strict contro
over environmental conditions and easy accessibility to manipulations but
suffer from lacking the native neuronal circuitry as it is found in vivo.
This major disadvantage can be at least partially circumvented by
utilizing organotypic brain slice cultures. Organotypic cultures allow
the investigation of delayed pathol. processes after hypoxic/hypoglycemic
insults and of the long-term effects of neuroprotective compds, in the
present report the authors describe the development of organotypic
hippocampal slice cultures maintained on membrane filter inserts at the
interface between tissue culture medium and atmospheric as an in vitro
l for

I for the investigation of neuroprotective drugs against ischemic damage. Ischemia was simulated in vitro by combined oxygen/glucose deprivation. Neuronal cell death as measured by propidium iodide uptake 24 h after the insult was compared with functional damage as estimated in the short-term range by electrophysiol. recordings of field potentials. Pharmacol. validation was achieved by testing the effects of cytoprotective compds. with different effector mechanisms. Bearing in mind that OSC prepared from neonate rats may not represent the situation found in the adult CNS, they provide an exptl. in vitro system that is well suited to complement in vivo prepns. and dissociated cell cultures in studying long-term pathophysiol. processes of neurodegenerative diseases.

4368-28-9, TTX

REL BAC (Blological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

[protection of organotypic hippocampal slice cultures from ischemic

injury) 4368-28-9 HCAPLUS

5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,

ANSWER 63 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 2000:74515 HCAPLUS MENT NUMBER: 133:589

DOCUMENT NUMBER:

TITLE:

133:589
Neuroprotection against ischemia by metabolic inhibition revisited: A comparison of hypothermia, a pharmacologic cocktail and magnesium plus mexiletine Maynard, Kenneth I., Quinones-Hinojosa, Alfredo; AUTHOR (S):

Malek, Junaid Y.

CORPORATE SOURCE: Neurophysiology Laboratory, Massachusetts General Hospital and Harvard Medical School, Boston, MA,

Hospital and Harvard medical school, busicul, no., 02114, USA
Annals of the New York Academy of Sciences (1999), 890 (Neuroprotective Agents), 240-254
CODEN: ANYAA9, ISSN: 0077-8923
New York Academy of Sciences SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

English UNGS: English
Previous studies have suggested that metabolic inhibition is
neuroprotective, but little evidence has been provided to support this
proposal. Using the in vitro rabbit retina preparation as an established

model

of the central nervous system (CNS), the authors measured the rate of
glucose utilization and lactate production, and the light-evoked compound
action

action

potentials (CAPs) as indexes of neuronal energy metabolism and
electrophysiol.

function, resp. The authors examined the effect of three (3) treatments
options: hypothermia (i.e., 33° and 30°), a six-member
pharmacol. "cocktail" (tetrodotoxin (0.1 µM), 2-amino-4phosphonobutyric acid (20 µM), 2-amino-5-phosphonovaleric acid (1 mM),
amiloride (1 mM), magnesium (10 mM) and lithium (10 mM)) and the
combination of magnesium (Mg2-1 mM) and mexiletine (Mex, 300 µM) on in
vitro rabbit retinas, to see if there is a correlation between neuronal
energy metabolism during ischemia (simulated by the reduction of oxygen
from 55%
to 15% and glucose from 6 mM to 1 mM), and the subsequent recovery of
function. Bypothermia and the "cocktail" significantly inhibited both the
rate of glucose utilization and lactate production, whereas Mg2+ and/or Mex
showed only a nonsignificant tendency toward a reduction, compared to

rol retinas. Recovery of light-evoked CAPs was significantly improved in hypothermia- and cocktail-treated retinas, as well as with retinas exposed to the combination of Mg2+ plus Nex, but not with Mg2+ or Mex alone, relative to control retinas. A linear regression anal. of the % recovery of function vs. the % reduction in the rate of glucose utilization during ischemia showed a significant correlation (r2 = 0.80, correlation [ricient =

Ischemia showed a significant correlation (r2 - 0.80, correlation ficient = 0.9) between these two parameters. This and other data discussed provide convincing evidence that there is a correlation between metabolic inhibition, achieved during ischemia, and neuroprotection. 4368-28-9, Tetrodotowin RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (neuroprotection against ischemia by metabolic inhibition revisited and a comparison of hypothermia and pharmacol. cocktail and magnesium plus mexiletine)
4368-28-9 HCAPUUS
5,9:7,102-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

L8 ANSWER 63 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

67 THERE ARE 67 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 65 OF 108 HCAPAUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
132:276201
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polarizing (control) solution Sequential adultion to the Solution Sup 200

I) of optimal concess of HDE694 (Group II), furosemide (Group III), and BDM (Group IV) were compared with STB2 (Group V); postischemic recovery of acrtic flow was 29478, 49468*, 56428*, 76438*, and 25468, resp. (*PCO.05 vs. I and V). Creatine kinase leakage was lowest, and myocardial AFP content was highest in Group IV. Conclusions: A polarizing preservation solution (RH+TTM) containing HDE694, furosemide, and BDM significantly enhanced long-term preservation compared with an optimized depolarizing solution (STB2) used clin. For long-term donor heart preservation.
4368-28-9, Tetrodotomin
RL: BAC (Biological activity or effector, except adverse): BSU (Biological

preservation.

4368-28-9. Tetrodotoxin

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TBU (Therepeatite use); BIOL (Biological study); USES (Uses)

(sodium channel inhibitor; beneficial effects of polarized arrest (Na+-channel blockade), Na+/RH-exchange inhibition and Na+/K+/2Cl--cotransport inhibition combined with calcium desensitization on long-term beart preservation)

4368-28-9 HCAPUIS

5,9:7,10a-0imethano-10aH-[1,3]diomocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hemahydro-12-(hydromymethyl)-,

(4R,4aR,5R,7s,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: Journal

English

NGUAGE: English
The mechanism underlying the intestinal fluid loss in rotavirus diarrhea, which often afflicts children in developing countries, is not known. One hypothesis is that the rotavirus evokes intestinal fluid and electrolyte secretion by activation of the nervous system in the intestinal wall, the enteric nervous system (ENS). 4 Different drugs that inhibit ENS functions were used to obtain exptl. evidence for this hypothesis in mice in vitro and in vivo. The involvement of the ENS in rotavirus diarrhea indicates potential sites of action for drugs in the treatment of the

indicates potential sites of action for drugs in the treatment of the disease.

4368-28-9. Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(enteric nervous system in the fluid and electrolyte secretion in cotavirus diarchea)

4368-28-9 HCAPLUS

5,9:7,10a-Dimethano-10aH-(1,3)dioxocino[6,5-d)pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,

(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 34

ANSWER 65 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

52 THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 66 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

ECAPLUS COPYRIGHT 2006 ACS on STN
1999:715350 HCAPLUS
131:317714
Morphine contracts the guinea pig ileal circulating
muscle by interfering with a nitric oxide mediated
tonic inhibition
Lenard, Laszlo, Jr.; Halmai, Vilnos; Bartho, Lorand
Dep. Pharmacology Pharmacotherapy, Medical School,
Univ. Pecs, Pecs, H-7643, Hung.
Digestion (1999), 60(6), 562-566
COUEN: DIGEBW; ISSN: 0012-2823
S. Karger AG
Journal AUTHOR (5): CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: Journal English

UAGE: English
The effect of morphine was examined on the circular muscle of guinea pig
ileal segments in vitro, with special regard to its interaction with
enteric NO releasing neurons. In the presence of atropine (10-6 M),
morphine (10-6 M) caused tonic contraction (approx. 7% of the maximal
spassm) which was reversed by naloxone (10-6M). Tetrodotoxin (TTX: 10-6 M
also caused contraction (14% of maximum); morphine completely lost its

also caused contraction (14% of maximum) morphine completely lost its ect in the presence of TTX. Likewise, the NO synthase inhibitor NC-nitro-L-Arg (L-NOANG, 10-4 M) elicited a tonic circular muscle contraction (12% of maximum) and completely prevented the excitatory action of TTX or morphine. The NO donor Na nitro prusside (10-7-10-4 M) caused relaxation. In longitudinally oriented prepna. in the presence of atropine (10-6 M), TTX (10-6 M), or L-NOANG (10-4 M). In the circular muscle in the absence of atropine, cholecystokinin octapeptide (CCK-8; 10-9 M) evoked a tonic-phasic contractile response which spontaneously faded away within 3 min. L-NOANG (10-4 M) failed to affect intensity or duration of the response to CCK-8. It is concluded that NO-releasing myenteric neurons exert a tonic inhibitory influence upon the circular, but not longitudinal muscle of the guines pig 1seum. Morphine and TTX probably contract the circular muscle by reducing the amount of NO released. A release of NO seems to play no role in the contractile effect of CCK-8 or in its spontaneous termination.

4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TRU (Theoremeutic use); BIOL (Biological study); USES (Uses)
(TTX and morphine effects on the iteal circulating muscle by interfering with a NO mediated tonic inhibition)

4368-28-9 HCARUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino(6,5-d)pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute Stereochemistry.

L8 ANSVER 67 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1999:551459 HCAPLUS 132:117410 TITLE: Tetrodopovia accession and the state of the state o Tetrodotomin prevents posttraumatic epileptogenesis in

TITLE: Tetrodotoxin prevents posttraumatic epileptogenesis in rats

AUTHOR(S): Graber, Kevin D.: Prince, David A.

Department of Neurology and Neurological Sciences, Stanford University Medical Center, Stanford, CA, 94305-5300, USA

Annals of Neurology (1999), 46(2), 234-242

CODEN: ANNEDS: ISSN: 0364-5134

Lippincott Williams & Wilkins

DOCUMENT TYPE: Journal

ANGUAGE: Lippincott Williams & Wilkins

Journal

Lingliacer & Bajlish

AB Severe cortical trauma frequently causes epilepsy that develops after a long latency. We hypothesized that plastic changes in excitability during this latent period might be initiated or sustained by the level of neuronal activity in the injured cortex. We therefore studied effects of action potential blockade by application of tetrodotomin (TTX) to areas of cortical injury in a model of chronic epileptogenesis. Partially isolated islands of sensorimotor cortex were made in 28- to 30-day-old male

Sprayue-Dawley rats and thin sheets of Elvax polymer containing TTX or control

Sprague-Davley rate and thin sneets on any payment of the control vehicle were implanted over lesions. Ten to 15 days later necocrtical slices were obtained through isolates for electrophysiol. Studies. Slices from all animals (n = 12) with lesions contacted by control-Elvax (58% of 36 slices) exhibited evoked epiteptiform field potentials, and those from 4 rate had spontaneous epileptiform events. Only 2 of 11 lesioned animals and 66 of slices from cortex exposed to TTX in vivo exhibited evoked epileptiform exentials, and no spontaneous epileptiform events were observed

epileptiform potentials, and no spontaneous epileptiform events were rived

There was no evidence of residual TTX during recordings. TTX-Elvam was ineffective in reversing epileptogenesis when implanted 11 days after cortical injury. These data suggest that development of anti-epileptogenic drugs for humans may be possible.

4368-28-9, Tetrodotomin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TMU (Therapeutic use); BIOL (Biological study); USES (Uses)

(tetrodotomin prevents posttraumatic epileptogenesis in rats)

4368-28-9 ECAPUS

5,9:7,10a-Dimethano-10aH-[1,3]diomocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hemahydro-12-(hydromymethyl)-,

(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 66 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 67 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 70 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 70

L8 ANSWER 68 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1999:465416 HCAPLUS
DOCUMENT NUMBER: 132:102575
TITLE: Effects of tetrodotoxin and OKY-046 in renal ischemia

reperfusion Garvin, Paul J.: Niehoff, Michael L.: Robinson, Sandra AUTHOR(S):

M. Department of Surgery, Abdominal Organ Transplant Division, St. Louis University Health Sciences Center, St. Louis, MO, 63110-0250, USA Journal of Surgical Research (1999), 85(2), 273-278 CODEN: JSGRA2: ISSN: 0022-4804 CORPORATE SOURCE:

SOURCE:

Academic Press PUBLI SHER:

DOCUMENT TYPE: LANGUAGE:

LISHER: CAUGARY 1998: UNCAT TYPE: JOURNAL PRESS.

UNENT TYPE: Journal GUIAGE: English Isohemic reperfusion injury (IRI) contributes significantly to posttransplant graft dysfunction. An emphasis, therefore, has been directed toward the identification of novel renoprotective agents. In this study, the renoprotective effect of tetrodotoxin (TTX) alone, or in combination with a thromboxane synthetase inhibitor (OKY-046), was investigated in a 60-min warm ischemia, 72-h reperfusion, IRI rodent model.Unilateral nephrectomized rats were treated with the test vehicle alone, 1, 2, or 4 mp/kg of TKT for 2 mg/kg of OKY-046 i.v., either 15 min pre- or postischemia, or 2 mg/kg TTX administered simultaneously with OKY-046 (2 mg/kg), following the ischemic interval. Baseline, 24, and 72-h mean plasma creatinine (Ct) and urea nitrogen (BUN) were compared Maximal renoprotection was demonstrated by significantly improved 72-h Ct and BUN levels with the 2 mg/kg of TTX or with 2 mg/kg of OKY-046, each administered after ischemia (ischemic control Cr = 8.01 ± 1.07 mg/dL vs TTX = 3.84 ± 0.80 mg/dL, P = 0.008; vs OKY-046 = 4.0 ± 1.5, P + 0.008; vs OKY-046 = 4.0 ± 1.5, P + 0.008; vs OKY-046 = 4.0 ± 1.5, P + 0.008; vs OKY-046 = 4.0 ± 1.5, P + 0.008; vs OKY-046 = 52.6 ± 22.5, P = 0.008). The combination therapy utilizing TTX with OKY-046 resulted in reduced animal survival, demonstrating no renoprotection as measured with the bischem. parameters. These results support the renoprotective effects of TTX in a severe, rodent IRI model. The exact mechanism of action, as well as the therapeutic potential of TTX in preservation/transplantation, varrants further study. (c) 1999 Academic Press.

A569-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study) unclassified); TBU (Therapeution)

A569-28-9, Tetrodotoxin
A669-28-9, Tetrodotoxin and OKY-046 in renal ischemia reperfusion)

4569-28-9, Tetrodotoxin and OKY-046 in renal ischemia reperfusion)

4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

LE ANSWER 69 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1999:130120 HCAPLUS
DOCUMENT NUMBER: 130:347278

TITLE: Vanilloid receptor agonists potentiate the in vivo
local anesthetic activity of percutaneously injected
site 1 sodium channel blockers

AUTHOR(S): Kohane, Daniel S.; Kuang, Yur Lu, Nu T.; Langer,
Roberts Strichartz, Gary R.; Berde, Charles B.
CORPORATE SOURCE: Department of Anesthesia, Children's Hospital, Boston,
M., USA

SOURCE: Anesthesiology (1999), 90(2), 524-534

CODDY: AMESAV: ISSN: 0003-3022

PUBLISHER: Lippincott Williams & Wilkins
DOCUMENT TYPE: Journal

LANGUAGE: English

AB Background: Capsaicin, the pungent ingredient in chili peppers, is a
vanilloid with noxious and analgesic effects that inhibits
tetrodotowin-resistant sodium currents. Because tetrodotoxin-resistant
currents are found primarily in small-diameter nociceptor afferents of the
peripheral nerves, their inhibition may lead to selective analgesia.
Therefore, the authors evaluated the interactions between tetrodotxin, a
site 1 sodium channel blocker, and capsaicin on nerve blockade in vivo.
Methods: Percutaneous sciatic nerve injections with 0 to 9.9 ml capsaicin,
0 to 120 MM tetrodotoxin, or both were administered to male
Sprague-Dawley rats. Thermal nociceptive and motor blockade were
measured. Data were expressed as medians with 25th and 75th percentiles.
Results: Capsaicin produced a transient increase in thermal latency with
no effect on motor strength. Tetrodotoxin reduced motor strength for a
longer duration than nociceptive and motor blockade were
measured. Data was synergistic, as evidenced by (1) supraadiditive
prolongation of both nociceptive and motor block, with the effect of
capsaicin reversed by the vanilloid antagonist capsazepine, and (2)
synergism in the frequency that rats achieved maximal block shown by
isobolog, anal. The combination of tetrodotoxin and capsaicin. Conclusions: Site 1
sodium channel blockers and vanilloids have synergistic effects on nerve
blockade in vivo. These interactions

Absolute stereochemistry.

ANSWER 68 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 69 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 70 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1999:81580 HCAPLUS
DOCUMENT NUMBER: 130:148705
Use of neurotoxin therapy for treatment of neurological-urological conditions and related disorders
SINVENTOR(S): Schmidt, Richard A.; Kaula, Norbert F.
University Technology Corporation, USA PCT Int. Appl., 19 pp.
CODEN: PIXKD2
DOCUMENT TYPE: Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9903483			19980715
		BG, BR, BY, CA, CH, CN,	
DK, EE, ES,	FI, GB, GE, GH,	HU, IL, IS, JP, KE, KG,	KP, KR, KZ,
LC, LK, LR,	LS, LT, LU, LV,	MD, MG, MK, MN, MW, MX,	NO, NZ, PL,
		SK, SL, TJ, TM, TR, TT,	UA, UG, US,
U2, VN, YU,			
		UG, ZW, AT, BE, CH, CY,	
FI, FR, GB,	GR, IE, IT, LU,	MC, NL, PT, SE, BF, BJ,	CF, CG, CI,
CM, GA, GN,	GW, ML, MR, NE,	SN, TD, TG	
CA 2296720 CA 2505930	AA 19990128	CA 1998-2296720 CA 1998-2505930	19980715
CA 2505930	AA 19990128	CA 1998-2505930	19980715
CA 2505933	AA 19990128	CA 1998-2505933	19980715
CA 2521392	AA 19990128	CA 1998-2521392	19980715
AU 9883007	A1 19990210	AU 1998-83007	19980715
AU 743085	B2 20020117	CA 1998-2505933 CA 1998-2521392 AU 1998-83007	
EP 1011695	A1 20000628	EP 1998-933345	19980715
R: AT, BE, CH,	DE, DK, ES, FR.	GB, GR, IT, LI, LU, NL,	SE, MC, PT.
10 DT			
JP 2001510163	T2 20010731	JP 2000-502781 CN 1998-809129 CN 2003-2003110471 EP 2004-19371	19980715
JP 3692033	B2 20050907		
CN 1135986	B 20040128	CN 1998-809129	19980715
CN 1480212	A 20040310	CN 2003-2003110471	19980715
EP 1475099	A1 20041110	EP 2004-19371	19980715
EP 1475099	B1 20051228		
R: AT. BE. CH.	OE. DK. ES. FR.	GB, GR, IT, LI, LU, NL,	SE. MC. PT.
IE, FI, CY		,,,	,,,
EP 1502601	A1 20050202	EP 2004-26167	19980715
R: AT, BE, CH,	DE, DK, ES, FR.	GB, GR, IT, LI, LU, NL,	SE. MC. PT.
TE. FI. CY			
AT 314085 US 6365164 US 2002025327	E 20060115	AT 2004-19371	19980715
US 6365164	B1 20020402	AT 2004-19371 US 2000-463040	20000117
US 2002025327	A1 20020228		20011015
US 6667041	B2 20031223		
US 2004180065	A1 20040916		20030904
US 6667041 US 2004180065 US 2004126380	A1 20040701		20031014
us 7001602	B2 20060221		
US 2004259788	A1 20041223	US 2003-745332	20031222
US 2005048084	A1 20050303	US 2003-685995 US 2003-745332 US 2004-778924 US 2004-778948 JP 2004-367500	20040213
US 2005049175	A1 20050303	US 2004-778948	20040213
JP 2005089478	A2 20050407	JP 2004-367500	20041220
2000000	20000407	0. L004-301300	_0041220

L8 ANSWER 71 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1998:761807 HCAPLUS
DOCUMENT NUMBER: 130:17253
LOCAL anesthetic formulations
Kohane, Oaniel S., Berde, Charles B.; Strichartz, Gary
R., Langer, Robert S.
OURCE: Children's Hedical Center Corporation, USA; Brigham and Women's Hospital, Inc.
PCT Int. Appl., 50 pp.
CODEN: PIXX02
DOCUMENT TYPE: Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KINO	DATE	APPLICATION NO.	DATE
WO 9851290	A2	19981119	₩O 1998-US9991	19980515
WO 9851290	A3	19990211		
W: AU, CA, JP				

W: AU, CA, UP
RV: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
PT, 5E
AU 9873990 A1 19981208 AU 1998-73890 19980515
US 6326020 B1 20011204 US 1998-79622 19980515
PRIORITY APPLIAL INFO.: US 1997-61633P P 19970516

W: AU, CA, JP

RY: AT, BE, CH, CT, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

AU 9973990

Al 19981208

Al 19981209

Bl 20011204

US 1999-79622

19980515

US 6326020

Bl 20011204

US 1999-76622

19980515

US 1997-46633P

P 19970516

US 1997-46761P

US 1997-46761P

US 1997-46761P

US 1997-46761P

US 1997-53462P

US 1997-53462P

US 1997-53462P

P 19970723

WO 1998-US9991

Combinations of naturally occurring site 1 sodium channel blockers, such as tetrodotoxin (TTK), saxitoxin (STK), decarbamoyl saxitoxin, and neosaxitoxin (referred to jointly herein as "toxins"), with other agents, have been developed to give long duration block with improved features, including safety and specificity. The duration of the block is greatly prolonged by combining a toxin with a local anesthetic, vasconstrictor, glucocorticoid, and/or adrenergic drugs, both a-agonists

(epinephrine, phenylephrine), P-blockers (propranolol), and mixed central-peripheral a-2 agonists (clonidine), or other agents. In another embodiment, the duration of nerve block from toxin can be greatly enhanced by the inclusion of amphiphilic or lipophilic solvents. The effectiveness of these compns. is enhanced by microencapsulation within polymeric carriers, preferably biodegradable synthetic polymeric carriers of the effective block can be obtained using combinations of toxin with vanilloids. TTK (0.2%) was combined with 50% bupivacaine and 0.05% dewamethasone in poly(glycolic acid-lactic acid) (65:35) microepheres.

The carrier fluid contained 1:100.000 epinephrine to reduce toxicity. The average duration of the effective block was 7 days.

2360-28-9, Tetrodotoxin

RL: BAC (Biological activity or effector, except adverse): BSU (Biological study), USES (USes)

(10021 anesthetic formulations)

4368-28-9 HCAPLUS

5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol.2-amino-1.4,4a,5,9,10-hemanydro-12-(hydroxymethy))-, (4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8	ANSWER 70 OF 108	HCAPLUS	COPYRIGHT	2006	ACS on STN	(Continued)
	JP 2005089479	A2	20050407	JP	2004-367501	20041220
	US 2005159337	A1	20050721	บร	2005-77895	20050311
PRIC	RITY APPLN. INFO.:			US	1997-52580P	P 19970715
				CA	1998-2296720	A3 19980715
				EP	1998-933345	A3 19980715
				JP	2000-502781	A3 19980715
				WO	1998-US14625	W 19980715
				บร	2000-463040	Al 20000117
				US	2001-978982	A2 20011015
				US	2003-685995	A2 20031014

AB

Methods are provided for treating neurol.-urol. conditions. A2 20031014 Methods are provided for treating neurol.-urol. conditions. This is accomplished by administration of at least one neurotoxin. 4368-28-9. Tetrodotoxin RL: BAC (Biological activity or effector, except adverse): BSU (Biological study, unclassified): THU (Therapeutic use): BIOL (Biological study): USES (Uses) (neurotoxins for treatment of neurol.-urol. conditions and related disorders)

alsorders)
4168-229-9 HCAPLUS
5,9:7,10a-01methano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 71 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 72 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

HCAPLUS COPYRIGHT 2006 ACS on STN
1998:705872 HCAPLUS
130:105205
Tetrodotoxin: anesthetic activity in the
de-epithelialized cornea
Schwartz, Daniel N. J. Duncan, Keith G.; Fields, Howard
L.; Jones, Matthew R.
Department of Ophthalmology, UCSF, San Francisco, CA,
94143, USA
Graefe's Archive for Clinical and Experimental
Ophthalmology (1998), 236(10), 790-794
CODEN: GACODL, ISSN: 0721-832X
Springer-Verlag AUTHOR (5):

CORPORATE SOURCE:

SOURCE:

PUBLI SHER: Springer-Verlag

DOCUMENT TYPE: LANGUAGE:

CODEN: GACODL, ISSN: 0721-832X

Springer-Verlag

UMENT TYPE: Journal

GUAGE: English

Background: Tetrodotoxin (TTX) binds with high affinity to sodium channels and produces local anesthesia. We investigated whether TTX is an effective, long-acting corneal anesthetic in rabbits. Methods: After mech debridement of the central corneal epithelium, topical TTX (1 mM, 0.1 mM, or 0.01 mM) was applied to one eye each of 18 New Zealand White rabbits. The fellow eye of each rabbit was treated with control vehicle. Blink response to a mech. stimulus was assessed. Blink response was also assessed every 3 h for 30 h in 6 rabbits treated with 1 mM TTX administered every 6 h. In a sep. group of 12 rabbits with central epithelial debridement, the rate of epithelial healing was compared between animals treated with topical 1.0 mM TTX and animals receiving no treatment. Results: After 4 h, eyes treated with 1.0 mM and 0.1 mM TTX were still partially smesthetic. By 8 h, the mean anesthesia score for 1.0 mM TTX was approaching normal. With multiple dosing, all six rabbit eyes remained anesthetic for the duration of the experiment There was no significant difference in the rate of re-epithelialization between eyes treated with TTX and untreated controls. There was no evidence of systemic or local toxicity from topical TTX. Conclusion: In a rabbit model, TTX is a long-acting topical anesthetic that retains its effectiveness when administered repeatedly over 24 h and does not inhibit epithelial healing. It may have application in management of pain after photorefractive keratectomy.

2368-28-9, Tetrodotoxin
RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TMU (Therapsulte use); BIOL (Biological study, unclassified); TMU (Therapsulte use); BIOL (Biological study); USES (Uses)

(tetrodotoxin anesthetic activity in the de-epithelialized cornea) 4368-28-9, Tetrodotoxin Anesthetic activity in the de-epithelialized cornea) 4368-28-9 HCAPLUS

Absolute stereochemistry.

L8 ANSWER 73 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1998:682103 HCAPLUS
DOCUMENT NUMBER: 129:286010
Hethod of anesthesia using a long-acting sodium channel blocker
Schwartz, Daniel M., Fields, Howard L.
TO Regents of the University of California, USA PCT Int. Appl., 51 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. DATE

Absolute stereochemistry.

ANSWER 72 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

29

REFERENCE COUNT:

THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 73 OF 108 HCAPLUS COPYRIGHT 2006 ACS OR STN (Continued)

L8 ANSWER 74 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

HCAPLUS COPYRIGHT 2006 ACS on STN
1998:541187 HCAPLUS
129:211637
Hechanism of relaxant effect of clonidine in isolated
bovine tracheal smooth muscle
Arimitsu, Masaahi, Mitsui-Saito, Minori; Sato, Koichi;
Ozaki, Hiroshi; Koga, Yoshihisar, Karaki, Hideaki
Department of Anesthesiology, Kinki University School
of Medicine, Osaka, Japan
Journal of Pharmacology and Experimental Therapeutics
(1998), 286(2), 681-687
CODEN: JPETAB; ISSN: 0022-3565
Williams & Wilkins
Journal AUTHOR (5):

CORPORATE SOURCE: SOURCE:

PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE:

MAGE: English
The relaxant effect of clouddine and the possible involvement of
imidazoline II receptors in bovine tracheal smooth muscle (BTSM) were
examined Clouddine caused concentration-dependent significant relaxation

ISM precontracted with 0.1 or 1 µM carbachol (CCh) but not in 72.7 mM KCI-induced contraction. The relaxation in CCh-contracted BTSM was inhibited by yohimbine (1 µM) and idazoxan (10 and 30 µM) but not by tetrodotoxin, indomethacin and other adrenoceptor antagonists. Oxymetazoline (0.1-100 µM) and phentolamine (0.1-100 µM) caused concentration-dependent relaxation, which was attenuated by idazoxan (10

Norepinephrine (0.1-100 µM) produced concentration-dependent relaxation, which

nas completely abolished by propranolol (10 μM) but not by yohimbine (1 μM). In fura-PE3/AM-loaded BTSM, CCh and 72.7 mM KCI increased intracellular calcium concentration ({Ca++|i) followed by contraction. The high

Intracellular Calcium concentration ((La++1) followed by contraction. The K+-induced increase in [Ca++] was not affected by clonidine. In CCh-stimulated BTSM, clonidine decreased [Ca++] and muscle force in parallel, whereas verapamil decreased [Ca++] in more strongly than muscle force. Clonidine (100 pM) inhibited the transient increase in [Ca++] induced by CCh but not by caffeine (20 pM) in Ca++-free solution Clonidine did not change the cAMP content in the presence of either 72.7 mM KCI or CCh. These results indicate that clonidine relaxes CCh-stimulated BTSM through the inhibition of CCh-induced increases in Ca++-influx, Ca++-release and intracellular signal transduction probably via imidazoline II receptors.

4369-28-9, Tetrodotoxin
RL: BAC [Sological activity or effector, except adverse); BSU (Biological study, unclassified); TMU (Therepeutic use); BIOL (Biological study, unclassified); TMU (Therepeutic use); BIOL (Biological study, unclassified); TMU (Therepeutic use); BIOL (Biological study); USES (Uses)

smooth muscle)
smooth muscle)
smooth muscle)
smooth muscle)
5,9:7,103—Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hashydro-12-(hydroxymethyl)-,
(4M,4aR,57,59,510,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 75 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1998:477704 HCAPLUS
129:239784
171TLE: 129:239784
AUTHOR(S): Kohane, Daniel S.; Yish, Jamier Lu, Nu T.; Langer, Robert: Strichartz, Gary R.; Berde, Charles B.
Harvard Medical School, Massachusetts General
Hospital, Children's Hospital, Brigham and Women's Hospital, Boston, MA, 02115, USA
Anesthesiology (1998), 89(1), 119-131
CODEN: ANESAV ISSN: 0003-3022
Lippincott-Raven Publishers
DOCUMENT TYPE: Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

ISHER: Lippincott-Raven Publishers
MENT TYPE: Ochrani
NUMGE: Regish
Highly potent toxins such as tetrodotoxin that block sodium channels with
great specificity have been studied for many years and can provide
prolonged blockade when coordinistered with wasoconstrictors or
conventional local anesthetics. Their utility has been constrained,
however, by systemic toxicity. The authors examined the efficacy of
tetrodotoxin with and without epinephrine or bupiwacaine for producing
prolonged-duration sciatic nerve blockade in the rat, and they assessed
the degree of concomitant toxicity. Rats received percutaneous sciatic
nerve blockade using tetrodotoxin with and without epinephrine or
bupiwacaine. A subset received s.c. injections at the nuchal midline.
Nocloeptive, proprioceptive, and motor blockade were quantified using
contralateral leg responses as controls for systemic effects.
Tetrodotoxin without epinephrine produced sciatic nerve blockade, but with
considerable toxicity at most EDs. Epinephrine reduced the median
effective concentration of tetrodotoxin for nocloeption from 37.6 to 11.5

effective concentration of tetrodotoxin for nociception from 37.6 to 11.5

and prolonged its duration, such that reversible blocks lasting >13 h were
achieved. Epinephrine reduced measures of systemic distribution and
increased the median LD of tetrodotoxin from 40 to 53.6 nmole/kg, thus
more than quadrupling the therapeutic index. Bupivacaine increased the
local anesthetic potency of tetrodotoxin, reduced its systemic toxicity,
and, when coinjected >.c., increased the median LD from 43.7 to 47.7

nmole/kg. The addition of epinephrine did not further improve the
effectiveness of the bupivacaine-tetrodotoxin combination. Combinations
of epinephrine or bupivacaine with tetrodotoxin or with other high-potency
toxins active on sodium channels should be examined for the potential to
provide clin. useful, prolonged nerve blockade.

4368-28-9. Hetrodotoxin
AL: ADV (Adverse effect, including toxicity): BAC (Biological activity or
effector, except adverse): BSU (Biological study, unclassified): TBD

(Therapeutic use): BIOL (Biological study): USES (USes)

(bupivacaine or epinephrine interaction with tetrodotoxin for prolonged
duration local anesthesia)

4368-28-9 HCAPUIS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 74 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 75 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 36

L8 ANSWER 76 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

HCAPLUS COPYRIGHT 2006 ACS on STN
1998:339592 HCAPLUS
129:62683
Protection against myocardial ischemic/reperfusion
injury by inhibitors of two separate pathways of Na-

AUTHOR (S): CORPORATE SOURCE:

entry
Eng, Stanley; Maddaford, Thane G.; Kardami, Elissavet;
Pierce, Grant N.
Division of Stroke and Vascular Diseases, Inst. of
Cardiovascular Sciences, St. Boniface General Hospital
Res. Centre, Winnipeg, MB, Can.
Journal of Molecular and Cellular Cardiology (1998),
30(4), 829-835
CODEN: JRCDAY; ISSN: 0022-2828
Academic Press Ltd.

SOURCE:

PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English

DIRECT TYPE:

JOURNAL

SURGE: English

Previous work has demonstrated that drugs in combination will have an additive protective effect in Langendorff-perfused hearts. During reperfusion following 30 min of ischemia, developed tension and resting tension were 34t3 and 16225s, resp., of pre-ischemic values in non-treated ischemic hearts. The administration of HOR-642 to inhibit Na+/He exchange increased active developed tension (NT) to SIB128 of pre-ischemic levels and decreased resting tension (NT) to Illi33 of pre-ischemic levels. The administration of tetrodotoxin (TTX) to block the Na+ Channel increased DT to 56:38 of the pre-ischemic level and reduced the Rt to 1.56:123 of the pre-ischemic level. Together, HOR-642 and TTX increased recovery of DT to 63:23 of pre-ischemic level levels and improved RT to 16:44 of pre-ischemic levels after 30 min of reperfusion. All drug treatment protocols significantly lowered the creatine phosphokinase activity measured in the coronary effluent in comparison to that observed in the non-treated hearts. These data demonstrate that inhibition of Na+ entry through either Na+-Ht exchange or the Na+ channel protects the heart from ischemic injury, but there is no addhl. benefit of blocking both routes of Na+ entry simultaneously. This suggests that a threshold level of Nai+ may be a critical factor in ischemic ardioprotection.

4366-28-9, Tetrodotowin RL: BAC (Biological activity or effector, except adverse); BSU (Biological study), USIS (Uses)

(protection against myocardial ischemic/reperfusion injury by inhibitors of two sen, mathway of Na+ entry)

IT

(protection against myocardial ischemic/reperfusion injury by inhibitors of two sep. pathways of Na+ entry) 8-28-9 RCAPLUS

4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4aR,5R,75,95,105,10aR,115,125)- (9GI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 77 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1998:119609 HCAPLUS DCCUMENT NUMBER: 128:122452
TITLE: Commondation

128:132452
Compositions containing tetrodotoxin for use as analgesics and in termination of drugs of abuse

INVENTOR (S): PATENT ASSIGNEE (S): SOURCE:

Wang, Weiguo Wang, Weiguo, Peop. Rep. China Paming Zhuanli Shenqing Gongkai Shuomingshu, 7 pp. CODEN: CNOKEY

Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1145225	Α	19970319	CN 1996-119454	19960924
CN 1072486	В	20011010		
DIODITE SERVED .			CH 1006 110464	10050004

CN 1072486 B 20011010

PRIORITY APPLM. INFO.:

CN 1996-119454 19960924

AB Compns. containing tetrodotowin and their use as analgesics and for termination of drups of abuse are claimed. An injection for pain in cancer patients at the terminal stage contained tetrodotowin [0.5-10.0 pg/1-20 mL] and acetic acid [BH 4-5].

IT 4368-28-9. Tetrodotowin BIOL (Biological study), USES (Uses) (compns. containing tetrodotowin for use as analgesics and in termination of drups of abuse)

RN 4568-28-9 ECAPHUS (OHI-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (SCI) (CA INDEX NAME)

ANSWER 76 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 78 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN CAPLUS COPYRIGHT 2006 ACS on STN 1997:640562 HCAPLUS 127:298748 Injectable therapy with botulinum toxin for control of muscle spasms and pain related to muscle spasms Acki, Kei Roger; Wheeler, Larry A.; Garst, Michael E. Allergan, USA PCT Int. Appl., 55 pp. CODEN: PIXXUZ ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 1997-533754 ES 1997-916168 US 1996-619780 WO 1997-US4643 A 19960320 W 19970320

A method for administration of botulinum toxin, includes the steps of (a) selecting at least one neuromuscular blocking agent having a duration of activity shorter than neuromuscular blocking agent having a duration of activity shorter than neuromuscular blocking agent having a duration of activity shorter than neuromuscular blocking activity of botulinum toxin, (b) selecting at least one muscle of a muscle group; (c) i.m. injecting the selected agent into the selected muscle; (d) observing muscle relaxation in both the selected muscle and other non-selected muscles in the muscle group to determine spill-over, muscle tone and balance; (e) repeating steps (b) - (d) until a final muscle selection is found; and (f) i.m. injecting botulinum toxin into the final muscle selection.

4368-28-9. Tetrodotoxin
Ri: THU (Therapeutix use); BIOL (Biological study); USES (Uses)
[Anjectable therapy with botulinum toxin for control of muscle spasms and pain related to muscle apasms)

4368-28-9 BCAPUS
5,9:7,103-01methano-10aR-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,

ANSWER 78 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 79 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 79 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1997:573035 HCAPLUS DOCUMENT NUMBER: 127:243149

DOCUMENT NUMBER: TITLE:

127:243149
Sodium channel modulators prevent oxygen and glucose deprivation injury and glutamate release in rat neocortical cultures
Probert, A. W.; Borosky, S.; Marcoux, F. W.; Taylor,

C. P. Parke-Davis Research Division, Department of Neurological and Neurodegenerative Diseases, Warner-Lambert Company, Ann Arbor, MI, 48105, USA Neuropharmacology (1997), 36(8), 1031-1038 CODEN: NEPHEW 15SN: 0028-3908 CORPORATE SOURCE:

SOURCE:

PUBLI SHER: Elsevier DOCUMENT TYPE: LANGUAGE:

AUTHOR (S):

LISHER: Elsevier

MEDIT TYPE: Journal

UMGE: English

Neocortical cultures were deprived of oxygen and glucose to model ischemic neuronal injury. The authors used a graded series of periods of oxygen and glucose deprivation, providing graded insults. Cell death was seasured by release of lactate dehydrogenase (LDH). One hundred and twenty to 240 min of deprivation caused graded increases in glutamate overflow. LDH release and 45ca influx. Curves of LDH release with respect to deprivation time were shifted to longer intervals by treatment with tetrodotoxin (TTX, 3, 30 or 300 mM), phenytoin (10, 30 or 100 mM), lidocaine (10, 30 or 100 mM) or the N-methyl-D-aspartate antagonist CT? (3(2-catboxypiperazine-4-yl)proyl-1-phosphonic acid, 3, 10, 30 or 100 mM).

Iidocaine (10, 30 or 100 mM) or the N-methyl-D-aspartate antagonist CT? (3(2-catboxypiperazine-4-yl)proyl-1-phosphonic acid, 3, 10, 30 or 100 mM).

Iidocaine (10, 30 or 100 mM) or the N-methyl-D-aspartate antagonist CT? (3(2-catboxypiperazine-4-yl)proyl-1-phosphonic acid, 3, 10, 30 or 100 mM).

Iidocaine (10, 30 or 100 mM) or the N-methyl-D-aspartate antagonist CT? (3(2-catboxypiperazine-4-yl)proyl-1-phosphonic acid, 3, 10, 30 or 100 mM).

Iidocaine (10, 30 or 100 mM) or the N-methyl-D-aspartate antagonist CT? (3(2-catboxypiperazine-4-yl)proyl-1-phosphonic acid, 3, 10, 30 or 100 mM).

Iidocaine (10, 30 or 100 mM) or the N-methyl-D-indicate that NH at the N-methyl-D-indicate that NH at the NH a

Absolute stereochemistry.

L8 ANSWER 80 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1997:407334 HCAPLUS
DOCUMENT NUMBER: 127:104106
Beneficial effects of dilazep on the palmitoyl-L-carnitine-induced derangements in isolated, perfused rat heart: comparison with

Hara, Akiyoshi; Arakawa, Johji; Hashizume, Hiroko;

AUTHOR (S):

nuixo, fasushi Department of Pharmacology, Asahikawa Medical College, Asahikawa, 078, Japan Japanese Journal of Pharmacology (1997), 74(2), 147-153 CORPORATE SOURCE:

SOURCE:

147-153 CODEN: JJPAAZ; ISSN: 0021-5198 Japanese Pharmacological Society Journal

AGE: English
The present study was carried out to determine the effect of dilazep,

LANGUAGE: English

AB The present study was carried out to determine the effect of dilazep, having an inhibitory effect on the Na+ channel, on the mech. dysfunction and metabolic derangements induced by palmitoyl-L-carnitine in isolated rat heat and to compare the effect of dilazep with that of tetrodotoxin, a specific inhibitor of the Na+ channel. Bat heart were perfused aerobically at a constant flow according to Langendorff's technique and paced elec. Palmitoyl-L-carnitine (5 µM) decreased the left ventricular developed pressure and increased the left extricular eveloped pressure and increased the left extricular end diastolic pressure (i.e., it produced mech. dysfunction), decreased the tissue level of APP and increased the tissue level of denomine monophosphate (i.e., it produced metabolic derangements). These mech. and metabolic alterations induced by palmitoyl-L-carnitine were attenuated by either dilazep (1 µM) or tetrodotoxin (3 µM). Neither dilazep nor tetrodotoxin modified the mech. function and energy metabolism of the normal (palmitoyl-L-carnitine-untreated) heart. These results suggest that inhibition of the Na+ channel with dilazep or tetrodotoxin is responsible, at each part, for attenuating the palmitoyl-L-carnitine-induced mech. (36e-22-9, Tetrodotoxin THU (Therapeutic use), BIOL (Biological study, unclassified), THU (Therapeutic use), BIOL (Biological study), USES (Uses)

(beneficial effects of dilazep on palmitoyl-L-carnitine-induced derangements in isolated perfused rat heart and comparison with tetrodotoxin in relation to sodium channel blockade and energy metabolism

boliam)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hemahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,10s,10aR,11s,125)- (9CI) (CA INDEX NAME)

ANSWER 80 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 81 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 5,9:7,10a-Dimethano-104H-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,7S,95,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 81 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1997:199533 HCAPLUS DOCUMENT NUMBER: 126:287924

AUTHOR (S):

CORPORATE SOURCE:

SOURCE.

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

ANSVER 81 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ISSION NUMBER: 1997:199533 RAPPLUS
EE: The protective action of tetrodotoxin and
(1)-kavain on anaerobic glycolysis, ATP content and
intracellular Na* and Ca2+ of anoxic brain vesicles
(6)-kavain on anaerobic glycolysis, ATP content and
intracellular Na* and Ca2+ of anoxic brain vesicles
(GRATE SOURCE: Ulm, 89081, Germany
Neuropharnacology (1997), Volume Date 1996, 35(12),
1743-1752
CODEN: NEPHBW; ISSN: 0028-3908
Elsevier
MENT TYPE: Journal
Emplish
Because recent reports point to Na* channel blockers as protective agents
directed against anoxia-induced neuronal damage including protection of
anaerobic glycolysis, the influences of tetrodotoxin (TTX) and
(1)-kavain on anoxic rat brain vesicles were investigated with respect
to lactate synthesis, vesicular ATP content and cytosolic free Na* and
Ca2+ (Na*)i, [Ca2+]i, both of the latter determined fluorometrically
cying
SBFF and FURA-2, resp. After anoxia, basal lactate production was increase

Carr (Na*)1, (Lear)1, source to the state of the state of

half life (<1/2) of 14.5 min, indicating that anaerobic glycolysis was insufficient to cover the energy demand of anoxic vesicles. Correspondingly, [Na+]: and [Ca2+]i increased persistently after anoxia by 22.1 nM Na+ and 274.9 nM Ca2+, determined 6.3 min after onset. An addnl. stimulation of vesicles with verattidine accelerated the drop of ATF (<1/2 - 5.1 min) and provoked a massive Na+ overload, which leveled off to 119 mM Na+ within a few minutes. Concomitantly, [Ca2+]i increased linearly with a rate of 355 nsol Ca2+//min. Despite the massive perturbation of ion homeostasis, lactate production was unaffected during

first 8 min of veratridine stimulation. However, complete inhibition of lactate synthesis took place 30 min after veratridine was added. The Natchannel blockers TTX and (i)-kavain, if applied before anoxia, preserved vesicular ATP content, diminished anoxia-induced increases in [Na+]i and [Ca2+]i and prevented both the veratridine-induced increases of [Na+]i and [Ca2+]i and the inhibition of lactate production The data cate

rate a considerable Na+ influx via voltage-dependent Na+ channels during anoxia, which speeds up the decline in ATP and provokes an increase in [Ca2+]i. A massive Na+ and Ca2+ overload induced by veratridine failed to influence lactate synthesis directly, but initiated its inhibition.

4360-28-9, Tetrodotoxin

4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(protective action of tetrodotoxin and kavain on anaerobic glycolysis and ATP content and intracellular Na+ and Ca2+ of anoxic brain

vesicles) 4368-28-9 HCAPLUS

L8 ANSWER 82 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1997:87693 HCAPLUS
DOCUMENT NUMBER: 126:126698
IZITLE: 266698
AUTHOR(5): 267605 ACS on STN
AUTHOR(5): 167603 HCAPLUS
Sawanobori, Tohrus Adaniya, Hitoshi; Hirano, Yuji;
Hiraoka, Masayasus AUTHOR (5):

Savanobori, Tohru Adaniya, Hitoshi; Hirano, Yu Hiraoka, Masayasu Department of Cardiovascular Diseases, Medical Research Institute, Tokyo Medical and Dental University, Tokyo, Japan Japanese Heart Journal (1996), 37(5), 709-718 CODEN: JHSJAR, ISSN: 0021-4869 Japanese Heart Journal Association CORPORATE SOURCE:

SOURCE:

PUBLI SHER:

MENT TYPE: Journal WAGE: English The effects of antiarrhythmic agents, including Class I and IV drugs and 3-10 mM Mg2+, on acontine-induced arrhythmias were examined by using a conventional microelectrode and patch-clamp method in Langendorff-perfused rabbit hearts and isolated guines pig ventricular myocytes. Intracoronary administration of 0.1 µM acontine induced polymorphic ventricular tachycardia (PVT) which continued for >60 min. Addition of acontine to ventricular myocytes caused a prolonged action potential duration (APD) and the appearance of sarly after-depolarization (EAD), together with the occurrence of an invared hump of the I-V curve acound -60 to -40 mV and increased outward current at pos. voltages. Addition of 10 µM tetrodoconin (TTX) and 25 mM Mg2+ restored acontine-induced PVT to sinus rhythm in Langendorff-perfused prepns. and also shortened the prolonged APD, demonstrating the abolition of EAD by acontine in ventricular myocytes. However, antiarrhythmic actions of Mg2+ and TTX in acontine-induced arrhythmia are to abolish EAD and shorten the prolonged APD by suppression of the invard Na+ current around -60 to -40 mV. 4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse) BSU (Biological study, unclassified), TMU (Therapeutic use), BIOL (Biological study), USSS (Uses)

(acontine-induced heart arrhythmias response to)

study); USES (Uses)
(acontitne-induced heart arrhythmias response to)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol; 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

L8 ANSVER 82 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 83 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 83 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1997:16863 HCAPLUS DOCUMENT NUMBER: 126:207394 TITLE: 126:207394 Altered Na+-channel function as a

126:207394
Altered Na+-channel function as an in vitro model of
the ischemic penumbra: action of lubeluzole and other
neuroprotective drugs
Ashton, David Willems, Roland; Wynants, Jozef; Van
Reempts, Jos: Marrannes, Roger; Clincke, Gilbert
Department of Neuropsychopharmacology, Janssen
Research Foundation, Turnhoutseweg 30, Beerse, 2340, CORPORATE SOURCE:

Belg.
Brain Research (1997), 745(1,2), 210-221
CODEN: BRREAP: ISSN: 0006-8993 SOURCE:

PUBLI SHER: Elsevier DOCUMENT TYPE: LANGUAGE:

AUTHOR(S):

ISSEE: Elsevier
MENT TYPE: Journal
UAGE: English
Veratridine blocks Na+-channel inactivation and causes a persistent
Na+-influx. Exposure of hippocampal slices to 10 µM veratridine led to
a failure of synaptic transmission, repetitive spreading depression
(SD)-like depolarizations of increasing duration, loss of Ca+-homeostasis,
a large reduction of membrane potential, spongious dema and metabolic
failure. Normalization of the amplitude of the neg. DC shift evoked by
high K+ ACSF 80 min after veratridine exposure was taken as the primary
endpoint for neuroprotection. Compds. whose mechanism of action includes
Na+-channel modulation were neuroprotective (ICSO-values in µM):
tetrodotoxin 0.017, verapamil 1.18, riluzole 1.95, lamotrigine 210,
and diphenylhydantoh 16.1. Both NMDA (M-R-801 and APH) and non-NMDA
(NBOX) excitatory amino acid antagonists were inactive, as were
NOS-synthesis inhibitors (nitro-L-arginien and L-NMAP). (Ca2-channel
blockers (cadmium, nimodipine), and a K+-channel blocker (TEA).
Lubeluzole significantly delayed the time before the slices became
epileptic, postponed the first SD-like depolarization, allowed the slices
to better recover their membrane potential after a larger number of SD-like
DC depolarizations, preserved Ca2+ and energy homeostasis, and prevented
the neurotoxic effects of veratridine (ICSO-value 0.54 µM). A
entration

the neurotoxic effects of veratridine (ICSO-value 0.54 µM). A concentration of lubeluzole, which was 40+ higher than its ICSO-value for neuroprotection against veratridine, had no effect on repetitive Na+-dependent action potentials induced by depolarizing current in normal ACSF. The ability of lubeluzole to prevent the pathol. consequences of excessive Na+-influx, without altering normal Na+-channel function may be of benefit in stroke.

IT 4360-28-9, Tetrodotoxin RL: BAC (Biological activity or effector, except adverse); BSU (Biological activity or effector, except adverse); BSU (Biological study, unclassified), TMU (Therapeutic use); BJOL (Biological study, unclassified); TMU (Therapeutic use); BJOL (Biological study, unclassified)

Absolute stereochemistry.

L8 ANSWER 84 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1996:347149 HCAPLUS DOCUMENT NUMBER: 125:148803

AUTHOR (S) :

CORPORATE SOURCE:

PLATEUS
125:48803
Prevention of coxygenation-induced arrhythmias in guinea pig papillary muscles
Hayashi, Hideharur Terada, Hajimer Katoh, Hidekir McDonald, T. F.
Photon Med. Res. Cent., Hamamatsu Univ. Sch. Med.,
Hamamatsu, 413-31, Japan
Journal of Cardiovascular Pharmacology (1996), 27(6),
816-823
CODEN: JCPCDT, ISSN. 0250

CODEN: JCPCDT; ISSN: 0160-2446 Lippincott-Raven

816-823
CODEN: JCPCDT; ISSN: 0160-2446
Lippincott-Raven
DOCUMENT TYPE: Lippincott-Raven
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Effects of various agents on recxygenation-induced arrhythmias, action
potentials, and tension of guinea pig papillary muscles were recorded to
investigate the site of action. Triggered activities due to delayed
afterdepolarizations (DADs) and aftercontractions were ellicited on
recxygenation after 60-min substrate-free hypoxia. Low extracellular Ca2+
(0.1 mM) abolished arrhythmias, and high Ca2+ (4.9 mM) increased the
amplitudes of DADs and aftercontractions. D-600 at a high concentration (20
µM) decreased the incidence of arrhythmias and decreased the recovery
of developed tension after recxygenation. Ryanodine (1 µM) abolished
aftercontractions and arrhythmias but did not affect the recovery of
developed tension. Tetrodotoxin (TTX 3 µM) and nicorandil (100 µM)
decreased the incidence of arrhythmias, but did not affect the recovery of
developed tension or the amplitudes of aftercontractions. TTX caused only
a slight decrease in Ca2+ transients in a fluo-3-loaded guinea pig
venticular myocyte. The Ca2+ entry through the Ca2+ channels apparently
synchronized Ca2+ release from the saccoplasmic reticulum, and D-600 at
the high concentration apparently decreased the incidence of arrhythmias.

and nicorandil decreased arrhythmias, probably by decreasing the Na+
current or by increasing the ATP-sensitive K+ current, resp.
4368-28-9, Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); THU (Therapeutic use); BIOL (Biological
study); USES (Uses).

(prevention of reoxygenation-induced arrhythmias in guinea pig
papillary muscles)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl)-,
(48,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

L8 ANSWER 84 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 85 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 85 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1996:93087 HCAPLUS DOCUMENT NUMBER: 124:194121

Selective suppression of in vitro electrographic seizures by low-dose tetrodotoxin: A novel TITLE:

anticonvulsant effect

Burack, Michelle A.; Stasheff, Steven F.; Wilson, AUTHOR(S):

Wilkie A. Wilkie A. Medical Center, Duke University, Durham, NC, 27710, USA CORPORATE SOURCE:

Epilepsy Research (1995), 22(2), 115-26 CODEN: EPIRE8; ISSN: 0920-1211 SOURCE:

Elsevier

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

JISHER: Elsewier
MENT TYPE: Journal

WAGE: English

Localized injections of 50 µM tetrodotoxin (TTX) in rat hippocampal

slices blocked stimulus train-evoked electrog, seizures (EGSs) for several

hours. Responses to single stimuli were minimally altered during TTX

block of the EGSs. This selective reduction of epileptiform activity could

result from general blockede of action potentials in an anatomically

distinct group of neurons in the slice. To test this hypothesis, we

systematically mapped TTX injection sites in the hippocampal slice, and

found that TTX injections that blocked EGSs were nearly always located in

or invaded CAZ's stratum radiatum and/or stratum lacunosum-moleculare. A

high degree of recurrent activity in this region contributes to both

epileptiform activity and responses to single stimuli; hence our selective

inhibition of EGSs suggests a more pharmacol. specific anticonvulsant

effect of TTX. Consistent with this hypothesis, we found that low concns.

of TTX (S. 10, or 20 mM) in the perfusion medium blocked EGSs without

decreasing the amplitude of extracellular responses to single stimuli.

Polysynaptic activity and/or antidromic firing may be particularly

vulnerable to TTX action on voltage-gated sodium channels, due to their

lower the safety factor for action potential propagation. Selective

of this activity may disrupt the abnormal neuronal activity underlying

of this activity may disrupt the abnormal neuronal activity underlying

EGSs. 4368-28-9, Tetrodotoxin

RE: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THO (Therapeutic use); BIOL (Biological study, unclassified); THO (Therapeutic use); BIOL (Biological study); USES (Uses)

[elective suppression of in vitro electrog. seizures by low-dose

(8616ctive suppression of in vitro electroy, selzures by 100-0050 tetrodotoxin)
4368-28-9 HCAPUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSVER 86 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:967255 HCAPLUS
124:807
Use of amino group-containing hydrogenated quinazoline compounds and derivatives thereof for the termination of drug dependence
INVENTOR(S): PATENT ASSIGNEE(S): Naning Maple Leaf Pharmaceutical Co., Ltd., Peop. Rep. China
SOURCE: COEN: PIXXOZ
DOCUMENT TYPE: Patent

Patent Chinese 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA:	ENT	NO.			KIN	D	DATE			APP	LICAT	ION:	NO.		п	ATE	
WO											1995-						
	w:	AM,	AΤ,	ΑU,	BB,	BG,	BR,	BY,	CA,	CH	, CN,	CZ.	DE.	DK.	EE.	ES.	FI.
											LK,						
											, RU,						
			UA														
	RW:	KE,	MW.	SD.	SZ.	UG.	AT.	BE.	CH.	DE	DK,	ES.	FR.	GB.	GR.	IE.	IT.
											CI,						
			TD,													,	,
AU	9518	881			A1		1995	1003		NU '	1995-	1889	1		1	9950	311
EP	7509	09			A1		1997	0102			1995-					9950	
EP	7509	09			B1		2002	1211					-		-		
	R:	BE.	DE.	FR.	GB												
JP	0951			,	T2		1997	1014		JP '	1995-	5237	SA .		1	9950	311
RU	2168	331					2001	0610			1996-					9950	
	5846				λ		1998				1996-					9960	
ORIT	APP	LN.	INFO	. :							1994-					9940	
											1995-					9950	

AB This invention relates to the use of amino group-containing hydrogenated quinazoline compds. and derivs. thereof, such as tetrodotoxin, for the termination of drug dependence in humans. Amino group-containing hydrogenated quinazoline compds. are administered s.c., i.m. or i.v. to subjects, and the said drugs are alkaloids and nitrogen-containing non-amino acid commds. such as opium, morphine, and heroin. The therapeutic amino group-containing hydrogenated quinazoline commds. are nonhabit-forming and fast-acting and show min. side effects.

IT 406-22-9D, Tetrodotoxin, derivs.
Ri. THU (Therapeutic use); BIOL (Biological study); USES (Uses) (use of amino group-containing hydrogenated quinazoline compds. and derivs.

thereof for the termination of drug dependence)
4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

ANSWER 86 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 87 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 87 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1395:765847 HCAPLUS
123:18028
Alleviation of contractile dysfunction in ischemic hearts by slowly inactivating Na+ current blockers
L8 Grand, B.; Vie, B.; Talmant, J. M.; Coraboeuf, E.;
John, G. W.

John, G. W. Div. Cardiovascular Diseases, Cent. Recherche Pierre Fabre, Castres, 81106, Fr. CORPORATE SOURCE:

American Journal of Physiology (1995), 269(2, Pt. 2), H533-H540 SOURCE:

CODEN: AJPHAP: ISSN: 0002-9513 American Physiological Society Journal PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

MENT TYPE: Journal MURGE: English English The authors hypothesized that the slowly inactivating component of Naturett, which is an integral part of the Natwindow current, is a major pathway for Nat loading during myocardial ischemia. The putative protective effects of tetrodotoxin (TTX) and R-56865, at concms. that selectively blocked the Natwindow current, as assessed by action potential plateau shortening without affecting maximum upstroke velocity (Ymax), were examined in isolated Langendorff-perfused guinea pig hearts subjected to 50 min of normothermic global ischemia and 60 min of reperfusion. In papillary muscles, TTX reduced action potential duration at 210 nM but reduced Vmax only at 21 μM. R-56865 (10 nM-10 μM) failed to affect Vmax but concentration dependently reduced on

on H-10 pM) failed to affect Vmax but concentration dependently reduced con potential duration. Ischemia-induced cardiac dysfunction, including increases in left ventricular end-diastolic pressure and lactate dehydrogenase and creatine phosphokinase release at reperfusion, was attenuated by TTX and R-56865 (0.1-320 nM). Ischemic contracture (increase in left ventricular end-diastolic pressure) was abolished by drug concens, as low as 1 nM, whereas higher concens. (>10 nM) of TTX and R-56865 were required to restore systolic function at reperfusion. TTX and R-56865 had lttle or no effect on hemodynamic variables. Evidence is provided of pronounced and direct cardioprotective effects of low concens. of R-56865 and TTX in cardiac muscle during ischemia. The results indicate that these drugs can selectively attenuate the Na* window current without affecting the fast peak of the Na* current and that the slow component of Na current may constitute a pathway of early Na* loading in the ischemic myocyte.

4368-28-9. Tetrodotoxin
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study; USES (Uses)

(alleviation of contractile dysfunction in ischemic hearts by slowly inactivating Na* current blockers)

4368-28-9 HCAPLUS

5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 88 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 1995:505711 HCAPLUS HENT NUMBER: 122:292076

DOCUMENT NUMBER: TITLE:

AUTHOR (5):

122:292076

Sodium channel blockers reduce oxygen-glucose deprivation-induced cortical neuronal injury when combined with glutamate receptor antagonists by the combined with glutamate receptor antagonists. Hynch, James J., III; Yu, Shan P., Canzoniero, Lorella M. T.; Sensi, Stefano L.; Choi, Dennis W. Department Neurology, Washington School Medicine, St. Louis, MO, USA
Journal of Pharmacology and Experimental Therapeutics (1995), 273(1), 554-60
CODEN: JPETAB; ISSN: 0022-3565
Williams & Wilkins
Journal

CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE:

(1995), 273(1), 554-60

CODEN: JPETAB: ISSN: 0022-3565

LISHER: Villiams & Vilkins

JOURNAL

JUACE: English

Blockers of voltage-gated Na+ channels can protect central neuronal axons
from hypoxic injury in vitro but have shown limited neuroprotective
effects on neurons, where substantial injury is mediated by glutamate
receptors. The authors explored the ability of several voltage-gated Na+
channel blockers to protect murine cultured cortical neurons from injury
induced by oxygen-glucose deprivation. Whole-cell recordings from neurons
revealed two types of Na+ currents activated by membrane depolarization:
one rapidly inactivating and the other noninactivating. Both currents
were blocked by tetrodocxin (TTX) and S,5-diphenyldyantonin (phenytoin).
Fluorescent imaging using the Na+-selective dys SBFI confirmed that TTX
attenuated the increase in intracellular free Na+ induced by
oxygen-glucose deprivation. Addition of TTX (I aN) but not phenytoin
(10-100 µM) produced a small and variable reduction in neuronal death
subsequent to oxygen-glucose deprivation for 40 to 50 min. Blockade of
glutamate neurotoxicity by combined addition of MX-801, 7-chlorokynurenate
and 6-cyano-7-nitroquinoxaline-2, 3-dione markedly reduced injury such that
prolonged deprivation times (75-100 min) were needed to induce widespread
neuronal death. In this setting of glutamate receptor blockade, addition of
MY-314 (I mM), quinidine (100 µM) or lorazinide (10 or 100
µM)-all further reduced neuronal death. Present results raise the
possibility that Na+ channel blockers may be useful in protecting gray
matter from hypoxic-ischemic injury, especially when combined with
anti-excitotoxic approaches.
4368-28-9, Tetrodotoxin
RL: BAC Biological activity or effector, except adverse); BSU (Biological
study), USES (Uses)
(Sodium channel blockers reduce oxygen-glucose deprivation-induced
cortical neuronal injury when combined with glutamate receptor
antagonists)
4368-28-9 HCAPLUS
5,917,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pen

L8 ANSWER 88 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 89 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 89 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 1995:292090 HCAPLUS MENT NUMBER: 122:71814 ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

122:71814
Neuroprotective effects of tetrodotoxin as a Na+
channel modulator and glutamate release inhibitor in
cultured rat cerebellar neurons and in gerbil global

AUTHOR (S):

Cultured rat Cerebellar neurons and in gerbil global brain ischemia
Lysko, Paul G.; Webb, Christine L.; Yue, Tian-Li; Gu, Juan-Li; Feuerstein, Giora
Cardiovascular Pharmacology, SmithKline Beecham
Pharmaceuticals, King of Prussia, PA, 19406-0939, USA
Stroke (1994), 25[12], 2476-82
CODEN: SJCCA7; ISSN: 0039-2499 CORPORATE SOURCE:

DOCUMENT TYPE: Journal

MENT TYPE:

Journal
SIMORE: English terodotoxin-sensitive ion channels in
Studies examining the role of tetrodotoxin-sensitive ion channels in
hypoxic-ischemic neuronal damage have concluded that sodium influx is an
important initiating event. The authors examined the neuroprotectant effect
of tetrodotoxin on both cultured cerebellar neurons and on CAI hippocampal
neurons of gerbils exposed to brain ischemia. The authors studied
neuroprotective mechanisms using cultured rat cerebellar granule cells
exposed to veratridine, which induced cytotoxicity, neurotransmitter
release, and calcium influx. Survival of gerbil CAI neurons was examined by
direct neuron counts 7 days after 6 min of global ischemia with
reperfusion. Tetrodotoxin protected cultured neurons in a dose-dependent
manner from veratridine-induced [3H] aspartate efflux that was sodium
dependent, only 25% calcium dependent, and was inhibited by tetrodotoxin
(inhibitory concentration [CSD]-60 nmol/L). Veratridine initiated AB

dependent, only 25% calcium dependent, and was inhibited by tetrodotoxin (inhibitory concentration [ICS0]=60 mono/t]. Veratridine initiated increases in intracellular calcium that were also reversed by tetrodotoxin (ICS0=63 mono/t], reversal was dependent on the sodium-calcium exchânger and the sodium-potassium pump. Neuroprotection of 90% (vs. vehicle) of gerbil CA1 hippocampal neurons was achieved by pretreatment with 2 ng of tetrodotoxin delivered three times intracerebroventricularly, without causing hypothermia. Sodium channel blockers like tetrodotoxin may have utility in treatment of ischemic neuronal injury by preventing excessive neuronal depolarizations, limiting excitotoxic glutamate release through reversal of the sodium-dependent glutamate transporter, preventing intracellular calcium overload, preserving cellular energy stores, and allowing recovery of ionic homeostasis through operation of the sodium-calcium exchanger.

IT 4368-28-9, Tetrodotoxin
RLE BAC (Biological activity or effector, except adverse), BSU (Biological study, unclassified), TRU (Therapeutic use), BIOL (Biological study), USES (Uses)
(neuroprotective effects of tetrodotoxin as Na+ channel modulator and glutamate release inhibitor in cultured rat cerebellar neurons and in gerbil global brain ischemia)

gerbil global brain ischemia)
4368-28-9 HCAPUUS
5,9:7,108-01imethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(RR,4a,S,7,5,9,5,105,10ax,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 90 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1994:692678 HCAPLUS DOCUMENT NUMBER: 121:292678 TITLE: Damage from 1

121:2920/8
Damage from oxygen and glucose deprivation in hippocampal slices is prevented by tetrodotoxin, lidocaine and phenytoin without blockade of action

AUTHOR (S): CORPORATE SOURCE:

Diocane and pnenytoin without blockage of action potentials Weber, Mark L., Taylor, Charles P. Department of Neuroscience Pharmacology, Parke-Davis Pharmaceutical Research, Division of Warner-Lambert Co., 2800 Plymouth Rd., Ann Arbor, MI, 48105, USA Brain Research (1994), 664 (1/2), 167-77 CODEN: BRREAP, ISSN: 0006-8993

SOURCE:

Brain Research (1994), 664(1/2), 167-77

CODEN: BRRARP; ISSN: 0006-8993

Elsevier

Journal

AB In vitro ischemia (VIV) was simulated with rat hippocampal slices in medium lacking D-glucose, equilibrated with 951 nitrogen, 5% carbon dioxide. Within 5-8 min, synaptic potentials disappeared and a DC neg. shift (5-15 mV) occurred. Prolonged application of 95% oxygen and D-glucose 12 min later did not allow synaptic potentials to recover. Slices pretreated with sodium channel blocking drugs allowed synaptic potentials to recover after IVI. Tetrodotoxin (TTX, 100-600 mM), the anticonvulsant phenytoin (5.0 to 100 mM) and the local nesthetic lidocaine (2.0 to 200 mM) each delayed or prevented neg. DC shifts from IVI. Histol. examination showed that drug treatments also prevented CA1 pyramidal cell damage from IVI. Neuroprotection occurred without blocking synaptic potentials or presynaptic fiber volleys, suggesting relevance for treatment of brain ischemia.

It 4366-28-9. Sidiological activity or effector, except adverse); BSU (Biological study), USES (Uses)

(damage from hippocampal ischemia is prevented by tetrodotoxin and lidocaine and phenytoin without blockade of action potentials)

RN 4368-28-9 RCAPLUS

NS 9,57,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-mino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (48,4aR,5s,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

L8 ANSWER 91 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:646337 HCAPLUS DOCUMENT NUMBER: 121:246337 TITLE: Methods for a comment of the comment of the

121:246337
Methods for treating neurodegenerative diseases and disorders using N-{2,6-disubstituted aromatic)-N'-pyridinyl ureas and other anticonvulsant compounds

compounds
Taylor, Charles Price, Jr.; Weber, Mark Lawrence
Warner-Lambert Co., USA
PCT Int. Appl., 31 pp.
CODEN: PIXXD2 INVENTOR (S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE A2 A3 19940901 19941222 WO 9418972 WO 9418972 WO 1994-U51788 19940217

9418972 A3 19941222
W: AU, CA, CZ, FI, HU, JP, KR, NO, NZ, RU, SK
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, HC, NL, PT, SE
6133299 A20001017 US 1993-23016 19930225
9462695 A1 19940914 AU 1994-62695 19940217
APPLN. INFO: US 1993-23016 A 19930225
WO 1994-US1788 W 19940217 PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

NRITY APPLM. INFO::

US 1993-23016 A 19930225
WO 1994-US1788 W 19940217
R SOURCE(S):

MARPAT 121:246337
Neucodegenerative diseases or disorders are treated by administering a therapeutically effective amount of a compound having anticonvulsant properties which bind to Na channels and modulate the channel without blocking the channel, to prevent irreversible neuronal damage from conditions similar to ischemia. Known N-{2,6-disubstituted phenyl)-N'-3-and 4-pyridinyl ureas and pharmaceutically acceptable acid addition salts thereof, e.g. N-(2-chloro-6-methylphenyl)-N'-4-pyridinyl urea, and known anticonvulsant compds., e.g. ralitoline, phenytoin, lamotrigine, tetrodotoxin, lidocadne, and carbamazepine, are used for treating neurodegenerative disorders, perinatal asphyxia, Alzheimer's disease, fluntington's disease, Parkinson's disease, and amyotrophic lateral sclerosis. Treatment with N-(2-chloro-6-methylphenyl)-N'-4-pyridinyl urea provided protection of hippocampal slices from irreversible loss of synaptic potentials after brief application of conditions that mimic ischemia in vitro. N-(2,6-dichlorophenyl)-N'-4-pyridinyl urea was prepared from 4-aninopyridine and 2,6-dichlorophenyl-N'-4-pyridinyl urea was prepared from 4-aninopyridine and 2,6-dichlorophenyl-

anticonvulsant

comvulsant compds.)
4368-28-9 KCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexabydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 92 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1990:132107 HCAPLUS
TITLE: Antischusker:

L8 ANSWER 92 OF 109 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1990:132107 HCAPLUS
DOCUMENT NUMBER: 1990:132107 HCAPLUS
TITLE: Antiacrhythmic properties of tetrodotoxin against occlusion-induced acrhythmias in the rat: a novel approach to the study of the antiacrhythmic effects of ventricular sodium channel blockade
AUTHOR(S): Abraham, Shlomor Beatch, Gregory N.; MacLed, Bernard
A.; Walker, Michael J. A.
CORPORATE SOURCE: Fac. Med., Univ. British Columbia, Vancouver, BC, V6T
1V5, Can.
SOURCE: Journal of Pharmacology and Experimental Therapeutics (1989), 251(3), 1166-73
CODEN: JPETAB; ISSN: 0022-3565
DOCUMENT TYPE: Journal
AB Blockade of ventricular sodium conductance (gNa) is believed to play an important role in the beneficial antiacrhythmic effects of class I antiacrhythmic agents. The present study was undertaken to examine the importance of ventricular gNa blockade by assessing the antiacrhythmic profile of tetrodotoxin (TTX), a selective sodium channel blocker. Expts. were performed in pentobarbital-anesthetized and artificially ventilated rats. Tvo doses of TTX were tested for antiacrhythmic activity and a high dose (TTM, 5) ps/(gc fo bolus + infusion of 50 pg/(g/h) which blocked only meuronal activity, and a high dose (TTM, 5) bps/(gc fo bolus + infusion of 50 pg/(g/h) which also produced signs of ventricular gNa blockade in normal hearts. To control for the decreases in blood pressure and heart rate caused by TTX, hexamethonium, nitroprusside and propranolol vere also used. Only TTX possessed antiacrhythmic activity in rats subjected to myocardial inchemia (produced by ligation of the left anterior descending oconeary actery). TTXh reduced dY/dt maximum of the action potential as well as action potential height, and concemitantly prolonged the P-R and QRS intervals of normal hearts. Apparently, drugs which produced hypotension, bradycardia and loss of autonomic function vere not antiacrhythmic. On the other hand, the marked antiacrhythmic activity of TTXh apparently of (TTXh apparently of TTXh

Absolute stereochemistry.

ANSWER 91 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSVER 93 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1989:S15199 HCAPLUS COPYRIGHT 2006 ACS ON STN 111:115199
TITLE: Preparation

111:15199
Preparation, testing, and formulation of heterocyclopyrroloquinazolines as antiarrhythmics Franke, Albrechtr Ostersehlt, Berndr Schlecker, Rainerr Rendenbach, Beatricer Von Philipsborn, Gerda BASF A.-G., Fed. Rep. Ger. Offen., 9 pp.
CODEN: GWXXEX INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:			
PATENT NO.	KIND DATE	APPLICATION NO.	DATE
DE 3730718	A1 19890323	DE 1987-3730718	19870912
JP 01071881	A2 19890316	JP 1988-222578	19880907
EP 307814	A2 19890322	EP 1988-114755	19880909
EP 307814	A3 19900808		
EP 307814	B1 19920408		
R: AT, BE, CH,	DE, ES, FR, GB,	IT, LI, NL	
AT 74605	E 19920415	AT 1988-114755	19880909
ES 2032317	73 19930201	ES 1988-114755	19880909
CA 1331608	A1 19940823	CA 1988-576970	19880909
US 5214047	A 19930525	US 1988-243469	19880912
PRIORITY APPLN. INFO.:		DE 1987-3730718	A 19870912
		EP 1988-114755	A 19880909
OTHER SOURCE(S):	CASREACT 111:115	199: MARPAT 111:115199	

The title compds. (I, R = H, halo, OH, NO2, amino, acylamino, Cl-4 alkory, alkyl, alkylsulfonic acid; A = (Cl-4 alkyl-substituted) Cl-4 alkylene; X = (Substituted) Ph, naphthyl, heterocyclyll were prepared A mixture of 4-chloro-1-(4-methylehenyl)let.nae-1-one, 2-(2-aminophenyl)-4,5-dihydroimidazole, NaI, and EtOH were treated with 12 N HCl and the mixture was refluxed for 3h. The solvent was removed and the residue was heated for 4 h at 120° to give 71% 2,3,5,6,7,8-hexahydro-5-(4-methylphenyl)imidazol[,2-c]eyrrolo[,1,2-a]quinazoline. I prolonged QT times in guinea pigs with ED20's of 0.25-1.5 mg/kg i.v. 122478-34-69
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); TMU (Thermpeutio

L8 ANSWER 94 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1987:568491 HCAPLUS
DOCUMENT NUMBER: 107:168491

TITLE: Do antiarthythmic drugs act on the site of abnormal impulse generation or act on the normal myocardium?
Hashimoto, Reitaro, Mitsuhashi, Harumir Akiyama,
Kentaro; Komori, Sadayoshi
CORPORATE SOURCE: Hashimoto, Yamanashi Med. Coll., Yamanashi,
409-36, Japan
Japansee Circulation Journal (1987), 51(2), 196-202
CODEN: OCTAR2; ISSN: 0047-1828
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Locally-induced digitalis arrhythmia was produced to study whether
antiarthythmic drugs suppress arrhythmia by directly acting on the
abhornal impulse generation or by suppressing Na channels of normal
myocardium to make it unresponsive to abnormal impulses. Dogs were
thoractomized and the anterior descending artery (ADA) was isolated and
an addhl. 10 mg every 20 min of ouabain was injected directly into the
ADA produced ventricular tachycardia originating from the digitalis
intoxication. Locally injected class | antiarrhythmic drugs, including
tetrodotomin, were effective in suppressing this arrhythmia. However,
when iv. applied lidocaine was prevented from reaching the ADA area,
lidocaine was not effective in suppressing this arrhythmia. However,
when iv. applied lidocaine was prevented from reaching the ADA area,
lidocaine was not effective in suppressing this arrhythmia. Byperestly,
class | drugs produce antiarrhythmic effect by directly suppressing the
digitalis damaged area, not by suppressing the normal myocardium.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological
Study), USES (Uses)
(antiarrhythmic activity of, abnormal impulse generation area vs.
normal myocardium as action site of)

N 4368-28-9 (PAPRUS
CN 5,9:7,10a-Dimethano-10aH-(1,3)dioxocino(6,5-d)pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4a.5,9,10-hexalydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7s,9s,10s,10aR,11s,12s)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 93 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) use), BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of, as antiacrhythmic)
122478-34-6 KCAPLUS
2H-Pyriaido(1,2-c)pyrrolo[1,2-a]quinazoline, 3,4,5a,6,7,8-hexahydro-5a-(2-thienyl)- (9CI) (CA INDEX NAME)

L8 ANSWER 95 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1197:18629 HCAPLUS
106:18629
4-Amino-6,7-dimethoxyquinazoline derivatives
Yokoyama, Keiichir Kato, Kojir Kitahara, Takumir Ono,
Hiroyasun Nishina, Takashir Kumakura, Mikior Awaya,
Akirar Nakano, Takuo
Mitsui Petrochemical Industries, Ltd., Japan; Mitsui
Pharmaceuticals, Inc.
Jpn. Kokai Tokkyo Koho, 56 pp.
CODEN: JORGAF
DOCUMENT TYPE:
LANGUAGE:
Japanese

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Japanese 2

PATENT NO.	KIND	DATE	AP	PLICATION NO.		DATE
JP 61140568	A2	19860627	JP	1984-263015		19841214
JP 05028709	B4	19930427				
US 4734418	A	19880329	บร	1985-805905		19851206
CA 1307786	A1	19920922	CA	1985-497106		19851206
EP 188094	A2	19860723	EP	1985-309049		19851212
EP 188094	A3	19871223				
EP 188094	B1	19920318				
R: DE, FR, GB,	IT					
HU 42479	A2	19870728	ΗU	1985-4783		19851213
HU 198481	В	19891030				
PRIORITY APPLN. INFO.:			JP	1984-263015	A	19841214
			JP	1985-194968	A	19850905
			JP	1985-204463	Ä	19850918
OTHER SOURCE(S):	CASRE	ACT 106:18629				

CASREACT 106:18629

The title compds. (I: R = heterocyclyl: Rl = H, HeO: l = 2, 3), useful as antihypertensives, were prepared Thus, a mixture of 4-amino-2-chloro-6,7-dimethoxyguinazoline and 5,6-dihydro-6-exhlyl-5-oxo-2-piperazinopyrido[4,3-d]pyrimidine in Me2CHCH2CH2OH containing Et3N was refluxed for 4 h to give

83% I (R = Qr Rl = Hr l = 2). I at 1 mg/kg p.o. lowered the blood pressure in spontaneously hypertensive rats. Tablets containing I were prepared 104955-95-79

104965-69-79
RL: BMC [Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as antihypertensive)
Pyrido[4,3-d]pyrimidin-5(GH)-one, 2-[4-(4-amino-6,7-dimethoxy-2-

ANSWER 95 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) quinazolinyl)hexahydro-1H-1,4-diazepin-1-yl]-6-ethyl- (9CI) (CA INDEX NAME)

ANSWER 96 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 104965-69-7 HCAPLUS Pyrido(4,3-d]pyriadidn-5(6H)-one, 2-[4-(4-amino-6,7-dimethoxy-2-quinazolinyl)hexahydro-1H-1,4-diazepin-1-yl]-6-ethyl- (9CI) (CA INDEX

L8 ANSWER 96 OF 108			2006 ACS on STN						
ACCESSION NUMBER:		608919 HCAI	LUS						
DOCUMENT NUMBER:	105:2								
TITLE:	Quinazoline derivatives and antihypertensive								
	prepa	rations cont	aining them						
INVENTOR (5):	Yokoy	ama, Keiichi	, Kato, Koji, Kit	ahara, Takumi, Ohno,					
	Hiroy	asu: Nishina	, Takashi: Awaya,	Akira: Nakano,					
	Takuo	Watanabe,	Kazuyuki; Saruta,	Sakae: Kumakura.					
	Mikio		-	•					
PATENT ASSIGNEE(S):	Mitsu:	i Petrochemi	cal Industries, L	td., Japan; Mitsui					
	Pharm.	aceuticals,	Inc.						
SOURCE:	Eur. 1	Pat. Appl.,	235 pp.						
	CODEN	EPXX DW	• •						
DOCUMENT TYPE:	Paten	t							
LANGUAGE:	Engli:	ah .							
FAMILY ACC. NUM. COUNT	: 2								
PATENT INFORMATION:									
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE					
EP 188094		19860723	EP 1985-309049	19851212					
EP 188094		19871223							
EP 188094		19920318							
R: DE, FR, G	B, IT								
JP 61140568		19860627	JP 1984-263015	19841214					
	B4	19930427							
JP 62056488	A2	19870312	JP 1985-194968	19850905					
JP 03071430		19911113							
JP 62067077	A2	19870326	JP 1985-204463	19850918					
JP 05029223	B4	19930428							
PRIORITY APPLN. INFO.:			JP 1984-263015						
			JP 1985-194968						
			JP 1985-204463						

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Piperazinyl- and homopiperazinylquinazolines I (Rl = H, HeO; R2, R3 = H, alkony; R4 = H, NH2; R5 = substituted 2-pyrimidinyl, 2-pyridinyl, 2-quinolinyl, fused pyrimidinyl; n = 2, 3) were prepared as antihypertensives. Thus, 4-bensyl-l-pherazinecarboxamidine sulfate was cyclocondensed with MeCOC(COZMe):CEOMe to give pyrimidinecarboxylate II. This was amidated with ECNH2 and cyclocondensed with DMF to give pyridopyrimidinone III, which was debenzylated and condensed with 4-amino-2-chloro-6,7-dimethoxyquinazoline to give piperazinylquinazoline IV. In rats 1 mg IV/kg orally reduced blood pressure 23.0% after 6 h, the effect lasting 24 h. Tablets were prepared each containing I 1, starch 60, microcrystn. cellulose 35, light silica 3, and Mg stearate 1 mg. 104965-69-79
RN: BAC (Biological activity or effector, except adverse); BSU (Biological study); PREP (Preparation); TBU (Therapsutica use); BIOL (Biological study); PREP (Preparation); USES (Uses)

CASREACT 105:208919; MARPAT 105:208919

OTHER SOURCE(S):

L8 ANSWER 97 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1985:142963 HCAPLUS
DOCUMENT NUMBER: 102:142963
ROle of tetrodotoxin-pensitive ion channels in
evolvement and cessation of cardiac arrhythmia due to
myocardial ischemia
AUTHOR(S): Rozenshtraukh, L. V.; Anyukhovskii, E. P.; Sharov, V.
G.

AUTHOR(S):

myocardial ischemia
Rozenshtraukh, L. V.; Anyukhovskii, E. P.; Sharov, V.
G.
CORPORATE SOURCE:

SOURCE:

SOURCE:

USSR Cardiol. Res. Cent., Moscow, USSR
Cardiol.: Int. Perspect., [Proc. Vorld Congr.], 9th
(1984), Meeting Date 1982, Volume 2, 955-70.

Editor(s): Chazov, E. I.; Smirnov, V. N.; Oganov, R.
G. Plenum: New York, N. Y.
CODEN: S3HTAB
Conference
English
AB Tetrodotoxin (I) {4366-28-9} showed antiarrhythmic activity in dogs with
arrhythmia induced by coronary artery ligation at 2 µg/kg i.v., as well
as in isolated hearts from dogs 24 h after coconary artery occlusion (i.e.
during the late stage of infarction) at 4 + 10-8 g/ml. I also
potentiated the antiarrhythmic activity of ethmozine [29560-58-5] and
mexiletine [31828-71-4] in vivo.

In 4368-28-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study); USES (Uses)
(antiarrhythmic activity of)
RN 4368-28-9 HCAPLUS

N 5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-mino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(48,4aR,5R,75,95,10s,10aR,115,125) (9CI) (CA INDEX NAME)

L8 ANSWER 98 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1985:4046 HCAPLUS
102:4046
AUTHOR(S): CORPORATE SOURCE: Podzuweit, T.; Binz, K. H.; Schaper, V.
Max-Planck-Inst. Physiol. Clin, Res., Bad Nauheim,
D-6350, Fed. Rep. Gec.
SOURCE: Recent Advances in Cardiac Arrhythmias (1983), 1, 1-8
CODEN: RACAEX; ISSN: 0951-807X

DOCUMENT TYPE: Journal English

UAGE: English
Ventricular arrhythmias were induced in the intact nonischemic pig heart,
by slow subepicardial infusion (10 µL/min) of agents known to increase
myocardial cAMP. Arrhythmias could be induced by infusing 1 of the
following agents, dissolved in 2.5 mM cacil2-150 mM NaCl: noradrenaline
(NA), adrenaline 10-5M each isoproterenoi-10-6 M NaCl-dibutyryl-cAMP,
N6-monobutyryl-cAMP, 5.10-2M each 8-Br-CAMP-5.10-2M together with Ro
7-2956-5.10-4M. In the presence of myocardial ischemia arrhythmias could
also be induced by infusing caffeine, theophylline-5.10-2M, histamine,
glucagon, or dopamine-10-3M each. Other agents precipitating archythmias

also be induced by intusing carreine, theophylines, 10-241 histamine, glucagon, or dopamine=10-3M each. Other agents precipitating archythmias ouabain-10-5M and aconitine-10-6M. Prolonged infusion of the latter resulted in ventricular fibrillation. The induction of ventricular tachycardia (VT) by NA infusion was facilitated by simultaneously infusing Ca2+. The NA-Ca2+-VT could be abolished by the resp. infusion of pindolol-10-6M; propanolol-10-4M; verapamil, D 600-10-4M each; MMCL2-5.10-4M; NiCl2, CoCl2-2.5.10-3M each; acetylcholine, butyrylcholine-10-4M each; catabamylcholine, methacholine-10-6M each; bethanechol-10-5M or muscarine-10-6M. Biochem. anal. showed that CAMP was increased at the NA-Ca2+-infusion site when arrhythmias ensued and that both β-blockers and choline esters prevent such accumulation of CAMP. Ouring VT induced by NA-Ca2+-infusion tachycardia was stopped within 10-30 s by occluding the coronary artery supplying the infusion area. This ischemic effect was readily reversed by coronary reperfusion. Infusion of NA-Ca2+ outside the ischemic area (anterior descending coronary artery ligated 2-thirds of the way from its origin) consistently precipitated ventricular fibrillation within 6 min after coronary artery ligation. Myocardial CAMP mediates the effects on heart rhythm of addrenergic overstimulation and muscarinic receptor activation by modulating the slow Ca2+ invard current, preferably by non-ischemic or reperfused myocardium. 4368-28-9

RL: BAC (Biological activity or effector, except adverse): BSU (Biological study, unclassified): TMU (Therapautic use): BIOL (Biological study): USES (Uses)

(antiarrhythmic activity of, CAMP in relation to)

4368-28-9 - Dalmethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pantol, 2-amino-1,4,4s,5,9,10-hexahydro-12-(hydroxymethyl)-,(4R,4aR,57,85,10,50,8,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

La ANSWER 99 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 194:17436 HCAPLUS
DOCUMENT NUMBER: 190:17436
TITLE: Comparative effects of fast- and slow-ion channel
blocking agents on reperfusion-induced arrhythmias in
the isolated perfused rat heart
Winslow, E.1 Marshall, R. J., Hope, F. G.
CORPORATE SOURCE: Sci. Dev. Group, Organon Lab. Ltd.,
Newhouse/Lanarkshire, MLI SSH, UK
Journal of Cardiovascular Pharmacology (1983), 5(6),
928-36
CODEN: JCPCOT; ISSN: 0160-2446
DOCUMENT TYPE: Journal of Eardiovascular Pharmacology (1983), 5(6),
28-36
CODEN: JCPCOT; ISSN: 0160-2446
DATE of English
AB The effects of bepridil [64706-54-3] (1-4 µM), a new antianginal
agent, on reperfusion-induced arrhythmias (RA) in the isolated perfused
rat heart were compared with those of tetrodotoxin [4368-28-9] (0 16-1.57
µM), verapamil [52-53-9] (0.5-2 µM), diltiazem [42399-41-7] (1-2
µM), nifedipine [21829-25-4] (0.02-0.2 µM) and nitrorptic concns,
neither infedipine nor nitrendipine reduced the incidence of RA, whereas
the other 4 agents did. Protection against RA does not appear to be
related to coronary vasodilatation or to a reduction in the degree of
ischemia
as assessed by lactate dehydrogenase release. However, neg. chronotropism
appears to be relevant in the mechanism of action of the Ca antagonists.
Substantial protection against RA by all active drugs was associated with PR
prolongation and (or) actioventricular block or suppression of sinus
rhythm. Thus, bradycardia may play an important role in the
antiacrhythmic action of bepridil, but the relative contributions made by
inhibition of the inward Ca and (or) Na currents remain unclear.

IT 4368-28-9
RL: BAC (Biological activity or effector, except adverse); BSU (Biological

4368-28-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TMU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antiarchythmic activity of)
4368-28-9 HCAPLUS

5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 98 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 99 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) L8 ANSWER 100 OF 108 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

HCAPLUS COPYRIGHT 2006 ACS on STN 1977:50802 HCAPLUS 86:50802 Preventing metastasis and primary tumor growth of H. Ep. Number 3 Shen, Ysung-Ying; Gitterman, Charles O. Merck and Co., Inc., USA U.S., 3 pp. CODEN: USXCMM Patent English 2

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 3991192 PRIORITY APPLN. INFO.: US 1975-600554 US 1974-467239 19761109 A 19750731 A2 19740506

1-Mercapto-5-hydroxy-6,7-tetramethylene-s-triazolo[3,4-b]pyrimidine (I) [61413-52-3] prevents in ovo metastasis of human epidermoid carcinoma and exhibits antitumor activity against primary human epidermoid carcinoma and other tumors, such as adenocarcinoma and sarcoma. Dosage units containing 100-500 mg I were recommended. 61413-52-3
RL: BAC (Biological activity or effector, except adverse): BSU (Biological study, unclassified): TWU (Therapeutic use): BIOL (Biological study): USES (Uses) (neoplasm inhibitor) [61413-52-3] HCAPLUS [1.2,4]Triazolo[4,3-a]quinazolin-5(1H)-one, 2,3,6,7,8,9-hexahydro-1-thioxo-(9CI) (CA INDEX NAME)

L8 ANSWER 101 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1976:516725 HCAPLUS
DOCUMENT NUMBER: 85:116725
The local anesthetic activity of tetrodotoxin alone and in combination with vasoconstrictors and local anesthetics
AUTHOR(S): Adams, H. Jack; Blair, Murray A., Jr., Takman, Bertil

CORPORATE SOURCE:

DOCUMENT TYPE: LANGUAGE:

Anesthetics

Adams, H. Jack, Blair, Murray R., Jr., Takman, Bertil H.

PORATE SOURCE:

Res. Dep., Astra Pharm. Prod., Inc., Framingham, MA, USA

IRCE:

Anesthesia & Analgesia (Baltimore, MD, United States)

(1976), SS(4), 568-73

CODEN: AACRAT; ISSN: 0003-2999

Journal

GUAGE:

Tetrodotoxin (TTA) [4368-28-9], alone and in cochination with various vasoconstrictors and local anesthetics, was evaluated for its ability to produce peripheral nerve blocks in the rat and central neural block in the cat and dog. High frequency and long duration of block were attained in sufficiently high concans of TTX were used, although latency was long and high dosage produced systemic toxicity. Prequency and mean duration of block could be increased and systemic toxicity. Preduced if TTX was administered with a wasoconstrictive agent. Conventional local anesthetics also enhanced the nerve-blocking activity of TTX. When appropriate concus. of TTX and local anesthetics were used, a high frequency of blocks characterized by short latency and long duration were demonstrated. Some indirect evidence that local anesthetics enhance TTX activity by reversibly increasing the permeability of various neural barriers to TTX is presented.

ASS-29-9

RLI BAC (Biological activity or effector. Assertices.

ALBO-1273
RIL BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); TBU (Therapeutic use); BIOL (Biological study); DSS (Uses)

(anesthetic activity of, local anesthetics and vasoconstrictors effect

on)
436-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 100 OF 108 HCAPLUS COPYRIGHT 2006 ACS ON STN

ANSWER 102 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 1972:429843 HCAPLUS
T7:29843 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE: Pharmacology of tetrodotoxin and saxitoxin

Kao, C. Y.
Downstate Med. Cent., State Univ. New York, Brooklyn, AUTHOR (S): CORPORATE SOURCE:

UNITED TO Proceedings (1972), 31(3), 1117-23 CODEN: FEPRA7: ISSN: 0014-9446 Journal: General Review SOURCE:

DOCUMENT TYPE:

MENT TYPE:

JOURNAL General Review

WIGGE:

A review with 35 refs. Tetrodotoxin (I) [4368-28-9] and samitoxin (II)

[35523-89-8] although chemical different, interfered with the early
transiently open ionic channel through which Na [7440-23-5] lons pass in
most common excitable membranes. A synthetic guandifinium compound bearing
partial structural similarity to I resembled I qual. In having some
selective actions on the spike-generating process of the frog sartorium
muscle. This qual. resemblance supported the idea that the guandifinium
modety was important for the actions of I. In whole animals, I and II
caused severe hypotension. II was a weaker hypotensive than I, and
produced a late pressor effect that was due to catechol amine secretion.
4368-28-9

RL: THU (Therapeutic use), BIOL (Biological study), USES (Uses)
(pharmacology of)

(pharmacology of) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156-6) (156

L8 ANSWER 103 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1971:110240 HCAPLUS
TITLE: 74:110240 HCAPLUS
HITTER: 86fect of saline infusion on the respective antiarrhythmic effects of inipramine and tetrodotomin against aconstitne-induced arrhythmias in the rat
AUTHOR(S): Lagier, Georges, Auclair, Marie C., Lechat, Paul
Inst. Phartmacol., Ec. Med., Paris, Fr.
SOURCE: 10:401, 10:9-19
CODEN: THERAP, ISSN: 0040-5957
JOURNAI TYPE: JOURNAI JOU

DOCUMENT TYPE: LANGUAGE:

CODEN: THERAP: ISSN: 0040-5957

JOHENT TYPE: Journal

GUAGE: French

For diagram(s), see printed CA Issue.

Infusion of hypertonic Nacl solns. (1.5-4.5%) in anesthetized,
artificially ventilated rats suppressed the antiarrhythmic effects of
tetradotoxin (I) against aconitine-induced arrhythmias, but had no such
effect on the antiarrhythmic activity of imprenine (II). It was
suggested that the antagonistic effect of Na+ with I occurred in the
myometrium, and that the absence of such an antagonism with II mayhave
been due to strong tissue binding of II.

4368-28-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study); USES (Uses)

(antiarrhythmic activity of, sodium chlorid.

(antiatrhythmic activity of, sodium chloride effect of) 58-28-9 HCAPLUS

4368-28-9 HCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(4M,4aR,5R,75,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 105 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1969:437457 HCAPLUS
TITLE: Pharmacologic effects of tetrodotoxin; cardiovascular and antiarcythaic activities
AUTHOR(S): Bennetain, Martin: Pharmacologic effects of tetrodotoxin; cardiovascular and antiarcythaic activities
SOURCE: Indiana Univ., Bloomington, IN, USA
(1968) 125 pp. Avail: 69-4728
From: Diss. Abstr. B 1969, 29(9), 3422
Dissertation
English

L8 ANSWER 104 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1970:475367 HCAPLUS DOCUMENT NUMBER: 73:75367

DOCUMENT NUMBER: TITLE:

73:73367 Suppression of the antiarrhythmic effect of tetrodotoxin against aconitine in rat by perfusion of hypertonic sodium chloride Lagier, Georges: Auclair, Marie C., Lechat, Paul Inst. Pharmacol., U.E.R. Biomed. Cordeliers, Paris, AUTHOR(S): CORPORATE SOURCE:

SOURCE:

Fr. Comptes Rendus des Seances de l'Academie des Sciences, Serie D: Sciences Naturelles (1970), 270(26), 3325-8 CODEN: CHDDAT; ISSN: 0567-655X

Journal

DOCUMENT TYPE: French

KGUAGE: French Perfusion of a hypertonic NaCl (15-451) solution into anesthetized, artificially ventilated rats suppressed the antiarrhythmic effect of tetrodotoxin against aconitine nitrate-induced cardiac arrhythmias. 'effect was due to Na since perfusion of hypertonic glucose solns. was devoid of this activity. 4368-28-9

4368-28-9
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(antiarrhythmic activity of, hypertonic sodium chloride antagonism of)
4368-28-9 HCAPLUS

4.06-29-9 HANUS 5,9:7,10a-Dimethus - 10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol. 2-mino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-, (RR,4a, 8R,7s,95,105,10aR,115,125)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSVER 106 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN SSION NUMBER: 1968:494968 HCAPLUS MENT NUMBER: 69:94968

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

69:94968
Comparative pharmacological actions of ciguatomin and tetrodotomin, a preliminary account Ogura, Yasumin Nara, Junkon Yoshida, Tamao Dep. Toxicol. Pharmacol., Chiba Univ., Chiba, Japan Toxicon (1968), 6(2), 131-40
CODEN: TOXIAG: ISSN: 0041-0101

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE: Journal English

UMDE: angular The pharmacol. actions of a MeOH-soluble extract of ciguatoxin from Lutjanus bohar were compared with those previously reported for crystalline

AB The pharmacol. actions of a newn-soluble extent of Layarana from bohar were compared with those previously reported for crystalline tetrodotoxin in crayfish, mice, and rats. In mice, there were no significant differences in median lethal dosages (LDSO) of i.p. (560 mg./kg.) or orally (530 mg./kg.) and instead ciquatoxin. The LDSO for intracaudally administered ciquatoxin was 29.3 mg./kg. in crayfish. In rats, injected ciquatoxin (10-30 mg./kg.) depressed blood pressure, and this was accompanied by respiratory failure. In mice, ciguatoxin did not show physostigaine-like action on electroencephalograms. Progressively higher doses of ciguatoxin (500-1000 mg./kg., i.p.) depressed heart rate, and evoked arrhythmia and bradycardia followed by death of rats. Ciguatoxin and tetrodotoxin produced similar toxic symptoms, but there seemed to be qual. pharmacol. differences between them.

IT 4368-28-9 HCAPLUS

N. 59:7, 10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(48,4a,58,75,75,10a-Dimethano-10aR-11,3)dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,4a,5,9,10-hexahydro-12-(hydroxymethyl)-,(48,4a,58,75,75,10a-Dimethano-10aR-1,5)dioxocino[6,5-d]pyrimidine-4,7,10,11,12-pentol, 2-amino-1,4,13,12,12,10-(CA INDEX NAME)

L8 ANSWER 107 OF 108 HEAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1968:409494 HEAPLUS
DOCUMENT NUMBER: 69:9494

AUTHOR(S): Ogura, Yasumi, Mori, Yoko
Dep. Toxicol. Pharmacol., Chiba Univ., Chiba, Japan
CORPORATE SOURCE: Dep. Toxicol. Pharmacol., Chiba Univ., Chiba, Japan
SOURCE: European Journal of Pharmacology (1968), 3(1), 58-67
CODEN: EJPHAZ, ISSN: 0014-2999

DOCUMENT TYPE: Journal
ARBUAGE: English
AB The local anesthetic actions of intradermally given tetrodotoxin,
anhydrotetrodotoxin (AHT), monoformylanhydrotetrodotoxin (HFAHT),
deoxytetrodotoxin (DDT), methoxytetrodotoxin (MOT), ethoxytetrodotoxin
(EDT), tetrodoaminotoxin (TAT), diacetylanhydrotetrodotoxin (DAAHT), and
tetrodonic acid were tested in mice (O.011-58.2 mg./kg.), guinea pips (7.5
+ 10-5-7.8 + 10-1mH) and rabbits, and on desheathed crayfish
abdominal nerve fibers (3 tme 10-7-3 + 10-1 mH) and compared with
the effects of procaine and dibucaine. Of all the compds, tested, the
crayfish nerve fibers were most sensitive to the anesthetic action of
tetrodotoxin. In alkaline solution tetrodotoxin was more effective in the
desheathed nerve preparation and in neutral solution it was more effective in

desheathed nerve. This suggests that the active form is the cationic form of tetrodotoxin and it penetrates nerve tissue more rapidly as its uncharged form. Heating a tetrodotoxin solution at alkaline pH greatly

of tetroocosin and at penetrates nerve and a state of the content of the content

L8 ANSWER 108 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STM
ACCESSION NUMBER: 1967:489145 HCAPLUS
COCUMENT NUMBER: 57:89145 HCAPLUS
CORPORATE SOURCE: Structure and activity in tetrodotoxin derivatives
Deguchi, Takehiko
Mcd. Lab. Pharmacol. Central Res. Lab., Sankyo Co.,
Tokyo, Japan
SOURCE: Japanes Journal of Pharmacology (1967), 17(2), 267-78
CODEN: JTPAAZ; ISSN: 0021-5198
DOCUMENT TYPE: Journal
LANGUAGE: Beglish
GI For diagram(s), see printed CA Issue.
AB Pharmacol. properties of compds. structurally related to the neurotoxin,
tetrodotoxin (I, R = CH), were studied. All the compds., including
tetrodotoxin (Gooxytetrodotoxin (I, R = H), methoxytetrodoxotin (I, R =
CMe), ethoxytetrodotoxin (I, R = OEL), tetrodaminotoxin (I, R = MEZ),
anhydrotetrodotoxin (II, R = CH), Il-monofoxylanhydrotetrodotoxin
formate (II, Rl = R, R = CHO), 5,11-disactylanhydrotetrodotoxin (II, R1
R2 = AC), and tetrodonic acid (IIII), showed similar pharmacol. properties
in symptomatology in mice, blood pressure and respiration expts. in cats,
and in tests on nerve conduction-blocking activity in the frog sciatic
nerve and on spasmolytic activity in the guinea pig ileum. A comparison
of the quent. differences in the pharmacol. activity of the compds. showed
that neurotoxin activity depended on the integrity of the hemilactal
structure and Off groups in positions 4 and 9 and either or both of these
in positions 6 and 11.

17 4368-28-9
RL: TRU (Thexapeutic use), BIOL (Biological study); USES (Uses)
(pharmacology of)

4368-28-9
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(pharmacology of)
4368-28-9 BCAPLUS
5,9:7,10a-Dimethano-10aH-[1,3]dioxocino[6,5-d]pyrimidine-4,7,10,11,12pentol, 2-amino-1,4,4,5,9,10-hexahydro-12-(hydroxymethyl)-,
(4R,4aR,5R,7S,9S,10S,10aR,11S,12S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 107 OF 108 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)